LIQUID PHASE ADSORPTION OF PURE, BINARY AND TERNARY SYSTEMS OF n-PARAFFINS ON LMS-5A

A Thesis Submitted
In Partial Fulfilment of the Requirements
for the Degree of
DOCTOR OF PHILOSOPHY

By
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to the
DEPARTMENT OF CHEMICAL ENGINEERING
INDIAN INSTITUTE OF TECHNOLOGY, KANPUR
MARCH, 1979

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SYNOPSIS

Separation of n-paraffins from petroleum fractions has been of great importance in petroleum industry. Molecular sieves are being extensively used as adsorbent in these separation processes. Since most earlier processes use vapor phase adsorption, enough information has been generated on this aspect. However, present trend is towards the use of liquid phase adsorption which has been investigated by few workers. In the present study, liquid phase adsorption isotherms of several normal paraffins on LMS-5A were measured in the temperature range 6-42°C. A mathematical model based on statistical thermodynamics was developed to predict multicomponent adsorption, using adsorption isotherms of pure

components. The model was validated by experimentally measuring binary and ternary data on the same hydrocarbons.

A glass adsorption cell was developed for the measurement of adsorption of pure and multicomponent hydrocarbon systems. n-Pentane, n-hexane, n-heptane and n-Octane were used for liquid phase adsorption at 6°, 18°, 30°, and 42°C. Benzene served as a diluent since its adsorption is \ negligibly small on LMS-5A.

The general trend of the isotherms for pure n-paraffins showed an initial increase in equilibrium loading (m moles adsorbed/g of zeolite) with increase in concentration followed by a relatively large range of concentration over which the equilibrium loading was nearly constant and then a further increase at higher concentrations. This upward trend in equilibrium loading at higher concentration was attributed to adsorption on external surfaces rather than intracrystalline adsorption. The effect of temperature on equilibrium loading was maximum for n-pentane and reduced for higher paraffins, being minimum for n-octane. For a particular temperature, the effect of concentration on equilibrium loading was most significant for n-pentane and decreased with increase in molecular size. At a given temperature and molar concentration, equilibrium loading for lower molecular weight paraffin was higher than that for larger molecules. This was explained in terms of steric effect which is more predominant for larger molecules.

Pure component adsorption data were represented by a statistical model with two parameters namely, slope of the isotherm at low concentration, α , and volume of the adsorbed molecule, β. The grand partition function of the system defined in terms of α and β was used to calculate the number of molecules adsorbed per cavity. The model accounted for adsorbate-adsorbent interaction and the reduction in free volume due to the finite size of the adsorbed molecules. Accuracy required by the model and nature of the adsorption isotherms did not permit direct measurement of the slope at low concentrations. Consistent and satisfactory values of the molecular volume of various n-paraffins were also not available. A least square minimization procedure, using Blind Search Technique, was employed to evaluate α and β from the experimental data. For all the n-paraffins studied, α was found to increase with decreasing temperature and increasing molecular weight. Molecular volume, as expected, monotonically increased with temperature. This model was then extended to predict the adsorption in binary and ternary systems using the values of α and β obtained for pure components.

Experimental measurements on 5 binary and 3 ternary systems of these n-paraffins were made at four different temperatures. The effect of temperature and concentration on separation factor, K, defined below was studied

$$K = \frac{y_L/x_L}{y_H/x_H}$$

where y and x are mole fractions in adsorbed and liquid phases and L and H are lighter and heavier components respectively. In all binary systems, lower molecular weight paraffins were preferentially adsorbed. The value of K, at the same temperature, decreased with increase in composition of preferentially adsorbed component. To make the separation factor independent of concentration it was redefined as

$$K_{r} = \frac{y_{L}/x_{L}}{y_{H}/(x_{H}+x_{B})}$$

where \mathbf{x}_{B} is the mole fraction of benzene in the liquid phase. Adsorption measurements on ternary systems showed that total hydrocarbon adsorbed increased with decrease in temperature.

The extended form of the model was then validated against the experimental measurements of the binary and ternary systems. For most of the data, the predicted values of the number of molecules adsorbed per cavity were within 10 per cent of the experimentally observed values. This prediction procedure can be extended to systems of practical importance like separation of n-paraffins from petroleum fractions, which will facilitate their design without extensive pilot plant studies.

NOMENCLATURE

а	activity of the sorbate, p/kT
С	average number of molecules adsorbed per cavity
^c cal	average number of molecules adsorbed per cavity (calculated)
c _{expt}	average number of molecules adsorbed per cavity (experimental)
E	sum of square of errors
$^{\mathrm{E}}$ min	minimum of sum of square of errors
G	grand partition function
K	separation factor
Kr	redefined separation factor
k	Boltzmann constant
M	number of cavities
m	maximum number of molecules adsorbed in a cavity
$\overline{\mathbb{N}}$	average number of molecules adsorbed
n	number of data points in a adsorption isotherm
P	vapor pressure at adsorption temperature
p	partial pressure
Q	canonical ensemble partition function of the system
q.	partition function of a cavity
ri	position vector
S	number of molecules adsorbed in a cavity
T	absolute temperature, OK
Us	potential energy for the sub system

```
volume of the cavity (= 776 \text{ Å}^3 for LMS-5A)
\nabla
       mole fraction in liquid phase
\mathbf{x}
       mole fraction in adsorbed phase
У
       configuration integral of a cavity
Z
Greek Symbols
       Henry's constant. (molecules)/(cavity)(torr)
α
       molecular volume, A3
β
       molecular constant
ε
       diameter of the molecule
σ
       chemical potential
17
Subscripts
        component A
Α
В
        component B
C
        component C
       higher molecular weight paraffin
\mathbf{H}
        number of molecules of C adsorbed in a cavity
h
       number of molecules of A adsorbed in a cavity
i
        number of molecules of B adsorbed in a cavity
j
```

lower molecular weight paraffin

L

CHAPTER 1

INTRODUCTION

Amorphous, gel type, alumino silicates have been used for years for water treatment. An extensive literature survey till 1930 on the use of natural minerals, such as green sand or glauconite, and natural zeolite water softeners is given by Shreve [2]. Although, zeolite minerals were first discovered and named in 1756 by Baron Cronstedt, first experiments on separation of mixtures using dehydrated zeolite mineral chabazite were performed by Barrer only in 1938 [1]. Synthesis of molecular sieve zeolite in 1946 can be termed as a revolution in chemical industry. The magnitude of interest shown by scientists and engineers is reflected in the fact that from the first industrial research in 1948, over 7000 papers have been published and 2000 U.S. patents obtained dealing with zeolite science and technology.

Due to their many unique properties, zeolites are being used in a variety of applications in chemical industry. Broadly speaking, zeolites are used as adsorbents for separation processes and as catalysts or catalyst supports. Separation processes using molecular sieves as adsorbents include separation of n-paraffins from various petroleum fractions, separation of p- and o-xylenes, oxygen enrichment of air, removing water from water-alcohol mixture, drying of

refrigerants and other chemicals, recovering radioactive ions from waste solutions, removing carbon dioxide and sulfur compounds from natural gas, separating hydrogen isotopes, removing sulfur dioxide from flue gases, etc. Since very large surface areas are available for adsorption, molecular sieve zeolites are also used as catalysts or catalyst supports. For example, in the catalytic cracking of gas oil, zeolites are much superior than the conventional silica-alumina catalysts.

Separation of n-paraffins from petroleum fractions has been of great interest industrially. Higher carbon number n-paraffins recovered from kerosene or gas oil are used in the manufacture of biodegradable detergents while lower ones are removed from gasoline to increase its octane number. These n-paraffins when separated further can be used as organic chemicals. n-paraffins can not be separated from petroleum fractions by other conventional adsorbents such as activated carbons, activated clays, inorganic gels, etc.. because these adsorbents do not possess an ordered crystal structure and consequently the pores are non-uniform. Development of molecular sieves (LMS-5A) has made it possible to recover n-paraffins from petroleum fractions. Molecular sieves have a high internal area available for adsorption with uniform pore opening which makes them size selective. The adsorption takes place inside the cavity and the component having a critical diameter less than the mouth opening of the cavity is adsorbed while larger molecules are excluded. The only other method of n-paraffin separatiom is by forming adducts with urea. The formation of the complex is due to the ability of urea to form a frame work containing channels within which the molecules of n-paraffins fit lengthwise. This framework can accommodate molecules of different lengths but not those of widely different cross sections, i.e., the cross sections of branched paraffins, cycloparaffins, and aromatics are too large to permit these hydrocarbons to form adducts with urea. However, separation of n-paraffins by molecular sieves is superior to urea adduct method because of higher selectivity and easier generative properties of molecular sieves.

All petroleum fractions are mixtures of n-paraffins, iso- and other branched chain paraffins, olefins, naphthenes, etc. Each class of these components consist of several different hydrocarbons and each of these adsorb on molecular sieves to a different degree. The odsorbina, n-paraffins on LMS-5A is one to two orders of magnitude higher than that of other hydrocarbons. To design separation systems it is, therefore, necessary to study the multicomponent adsorption of these hydrocarbon mixtures on molecular sieves. Depending on the boiling range and origin, the composition of a petroleum fraction can vary widely and therefore there will be an

infinite number of multicomponent systems to be studied. It will be impossible to measure multicomponent adsorption data for each of these systems. However, the number of n-paraffins present in any petroleum fraction is small and adsorption studies can easily be made for each of these hydrocarbons. A model which could predict multicomponent adsorption using only pure component data would eliminate the need for multicomponent adsorption measurements.

The adsorption of n-paraffins can be performed in the vapor or liquid phase. Although vapor phase adsorption has been investigated by various workers, very little information is available on liquid phase adsorption. A systematic study on the liquid phase adsorption of n-paraffins on molecular sieves is very much needed.

Objectives of the Present Investigation:

In the present investigation, liquid phase adsorption studies on molecular sieves for the separation of n-paraffins from naphtha were envisaged. Since naphtha is a very complex mixture of hydrocarbons it was felt necessary to make rather simple mixtures of known components (n-pentane, n-hexane, n-heptane, n-octane and benzene) for this study. Benzene was used as a diluent since it is essentially not adsorbed on Linde Molecular Sieve 5A which was used as the adsorbent. Adsorption isotherms for these n-paraffins were measured at

several temperatures. These isotherms were then used in a mathematical model that was developed to predict the binary and ternary adsorption data. These predictions were verified by making actual measurements.

An overall literature review has been given in Chapter 2. Chapter 3 deals with the pure component liquid phase adsorption of these n-paraffins on molecular sieves. Experimental results of binary and ternary systems of these n-paraffins are given in Chapter 4. In Chapter 5 a mathematical model is described. Conclusions and recommendations have been discussed in Chapter 6.

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CHAPTER 2

LITERATURE REVIEW

2.1 HISTORICAL:

The phenomenon of adsorption was first discovered by Sheele and Fontana in 1777. The ability of charcoal to remove color from solutions was discovered by Lowtiz in 1785. deSaussure carried out the first systematic investigation of adsorption in 1814. The adsorption studies till 1930 have been reviewed by McBain [69]. A comprehensive review of physical adsorption was published in 1942 by Brunauer [26]. Later, Mantell reviewed adsorption as a unit operation of chemical engineering in 1951 [67].

The aluminosilicates were termed zeolites by Cronstedt [33] in 1756. He observed that certain mineral crystals when heated appeared to melt and boil at the same time. Thus, from the Greak Zeo, to boil, and lithos, stone, he coined the term zeolite. The phenomenon of melting and boiling at the same temperature is termed 'intumescence' and the zeolites are said to intumesce.

Initial investigations on crystalline zeolites were performed in 1840 by Damour [34] who reported on the reversible dehydration of the zeolite minerals. Damour noted that the transparency and crystal form did not change on heating.

In 1845, Thompson [98] conducted experiments which showed that certain soils have the power of decomposing and retaining ammonium salts. When a solution of ammonium sulfate was filtered through the soil, the filtrate contained calcium sulfate and ammonium salt was retained in the soil. Later, Way [101] showed that the hydrated silicates in the soil produced this phenomenon. He was able to show that only ammonium or potassium was exchanged for calcium in the soil. Several years later Eichhorn [40] in 1858 published a paper on the action of dilute salt solutions on silicates, showing that the base exchange principle discovered by Way is reversible. He studied the quantitative behavior of the zeolite minerals, chabazite and natrolite, in contact with dilute salt solutions and found that sodium and calcium could reversibly replace each other in these zeolites.

The idea that the structures of dehydrated zeolite consists of open sponge frame works, was given by Friedel [49] in 1896. He observed that various liquids such as alcohol, chloroform, carbondisulfide and benzene were occluded by zeolites. In 1909 Grandjean [53] found that dehydrated zeolite crystals could reversibly adsorb inorganic vapors such as iodine, mercury, ammonia, air, hydrogen, carbondisulfide and bromine.

Beginning in the 1920's, a large number of papers on crystalline zeolites began appearing in the scientific

literature. In 1925, Weigel and Steinhoff [104] reported that chabazite, rapidly adsorbed the vapors of water, methyl and ethyl alcohol and formic acid while acetone, ether, and benzene were excluded. This was the first report of the sieving action at molecular scale of anhydrous crystalline zeolites. Shortly, thereafter, McBain [69] recognized the importance of these results and deduced that the pore openings in the chabazite crystals must be less than 5 Å in diameter. To describe this phenomenon of selective adsorption, or 'persorption', as he termed it, McBain originated the term molecular sieve.

An extensive literature on the use of natural materials such as green sand or glauconite and zeolite water softners has been reported by Shreve [90]. Smith [93] in 1963 has defined zeolite as 'an aluminosilicate with a framework structure enclosing cavities occupied by large ions and water molecules, both of which have considerable freedom of movement, permitting ion exchange and reversible dehydration.'

The application of x-ray diffraction techniques by Pauling [81] and Taylor [97] in the early 1930's led to the determination of the crystal structures of analcite and natrolite. The porous nature of zeolite crystals attracted the attention of a few physical chemists, in particular Barrer, who reported the results of his studies on the sorption of gases on crystals of chabazite and analcite [8].

In the beginning, attempts [75] were made to duplicate the hydrothermal processes by which zeolite minerals were assumed to have formed in nature. Earlier scientists attempted to synthesize zeolite crystals with partial success. Although the synthesis of many zeolites has been reported in literature, majority of these must be discredited on the basis of improper identification [9-11]. The advent of x-ray diffraction has enabled a more positive identification of complex compositions and structures than the methods used by the early investigators. By 1952, many different types of synthetic zeolites had been prepared in the laboratory [9, 10, 18, 20, 67]. Some of these were new varieties not found in nature, e.g., type A and X. Recently, Bajpai [6] synthesized mordenite starting from rice husk ash which is available in abundance in India. Different zeolites were produced by changing the chemical composition and molecular weight distribution of the starting species in the silicate solutions. A selected list of natural and synthetic zeolites with their properties is given in Table 2.1.

2.2 STRUCTURE OF ZEOLITES:

2.2.1 Introduction:

Crystalline zeolites are complex materials, chemically and structurally, comprising the major group of the framework silicates [17]. Other minerals included in this category

TABLE 2.1: SOME ZEOLITES WITH THEIR PROPERTIES

Name	Source	Composition	Density, Void g/cm^3 cm	, .	Vol., Aperture,
alcire Group					:
Analcime	Natural	$\text{Na}_{16}[(\text{Alo}_2)_{16}(\text{Sio}_2)_{32}] \cdot \text{16} \text{ H}_20$	1.85	660.0	2.8
dalite Group		1			
Zeolite A	Synthetic	$\text{Na}_{12}[(\text{Alo}_2)_{12}(\text{Sio}_2)_{12}] \cdot 27\text{H}_20$	1,33	0*300	4.2
Faujasite	Natural	(Na ₂ , Ca, Mg) ₆₀ [(AlO ₂) ₆₀ (SiO ₂) ₁₃₂].			
		260H ₂ 0	1.31	0.350	8.0
Zeolite X	Synthetic	Nage[(AlO ₂)g6(SiO ₂) ₁₀₆].264 H ₂ 0	1.29	0.360	8 0
Zeolite Y	Synthetic	Na ₅₆ [(AlO ₂) ₅₆ (SlO ₂) ₁₃₆]·264 H ₂ 0	1.30	0.350	8.0
abazite Group					
Chabazite	Natural	$ca_4[(Alo_2)_8(Sio_2)_{16}] \cdot 26H_20$	1.48	0.290	3.0x4.3
Erionite	Natural	$(Ca, Mg, Na_2, K_2)_{45}[(Alo_2)_9(Sio_2)_{27}].$			
		27H ₂ 0	1.50	0,210	3.6x5.2
ilipsite Group		1			
Philipsite	Natural	$(K, Na)_5[(A10_2)_5(Si0_2)_{11}] \cdot 10H_20$	I	1	
Zeolite B	Synthetic	Na ₆ [(AlO ₂) ₆ (SiO ₂) ₁₀]·15 H ₂ 0	1.47	0.150	3.5

11

rdenite Group

Name

Mordenite

Mordenite

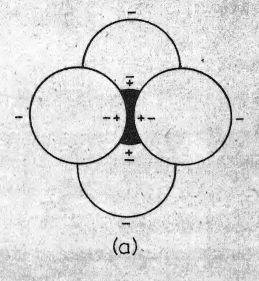
Source	Composition	Density, g/cm ³	Void Vol., cm ³ /g	Aperture,
	1			1
Natural	$^{\mathrm{Na}_{\mathrm{g}}[(\mathrm{AlO}_{2})_{\mathrm{g}}(\mathrm{SiO}_{2})_{40}]\cdot24\mathrm{H}_{2}^{\mathrm{O}}}$	1.72	0.150	4.0
Synthetic	$^{\text{Na}_{8}[(A10_{2})_{8}(Si0_{2})_{40}] \cdot 24_{\text{H}_{2}0}}$	1.72	0.140	9*9

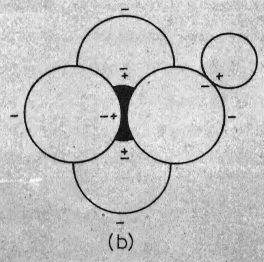
are feldspars and feldspathoids [36]. As a result of structural studies in the last three decades, using infrared, NMR and ESR in addition to x-ray diffraction, there is extensive information available on the structure of zeolites. Many zeolite properties can now be interpreted on the basis of their structure.

The first structural analysis of a synthetic zeolite, type A, was reported in 1956 [85] and was followed by the structures of faujasite [14], zeolite type X [19, 25] and chabazite [37]. These zeolites consist of a three dimensional frame work of SiO₄ and AlO₄ tetrahedra (Figure 2.1). Substitution of Al³⁺ for Si⁴⁺ in framework silicates is common. However, in order to maintain electrical neutrality, each substitution requires the presence of an alkali metal or alkaline earth ion, such as Na⁺, K⁺, Ca²⁺, Sr²⁺. In zeolites, the maximum substitution of Al³⁺ for Si⁴⁺ in case of A type zeolite is in the ratio of 1:1 and leads to a complete ordering of Al and Si ions [85]. The minimum substitution is in the ratio of 1:5 in case of mordenite [70]. One unit cell of the zeolite structure is given by

$$Me_{x/n} [(AlO_2)_x (SiO_2)_y] M \cdot H_2O$$

where Me is the metal cation, x and y are constants, n is the valency of the metal cation and M is the number of water molecules. The bracketted term gives the frame work composition.





ig. 2.1-Tetrahedra of four oxygen ions surrounding a silicon or aluminum ion. (a) Silicon-oxygen tetrahedra. (b) Aluminum-oxygen tetrahedra with Na cation.



Fig. 2-2 - Truncated octahedron.

The ratio of Al to Si, as discussed earlier, varies from l to 5.

The structure of many zeolites consists of simple arrangements of polyhedra; each polyhedra itself is a three dimensional array of Si AlO₄ tetrahedra in a definite geometric form. The sodalite group of zeolites (A, x and y types) are all based on frame works which are simple arrangements of truncated octahedra which contain eight hexagonal faces, six square faces, twenty four vertices and 36 edges [33] as shown in Figure 2.2. These truncated octahedra when interconnected in different ways form unit cells of different types of zeolites (A, x and y types).

2.2.2 Structure of Molecular Sieves 5A:

In the structure of zeolite type A [85] as shown in Figure 2.3, the octahedra are linked in cubic array by joining them with cubes on the square faces. This produces a central truncated cube octahedron with an internal cavity of 11 Å in diameter as shown in Figure 2.4. Each central cavity, termed the α cage, is entered through six circular apertures formed by a nearly regular ring of eight oxygen atoms with a free diameter of 4.2 Å. The cavities are, thus, arranged in a continuous three-dimensional pattern forming a system of unduloid-like channels with a maximum diameter of 11 Å and a minimum of 4.2 Å. The truncated octahedra

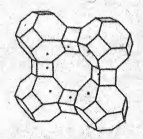


Fig. 2.3 - Cubic array of truncated octahedra in the zeolite type A.

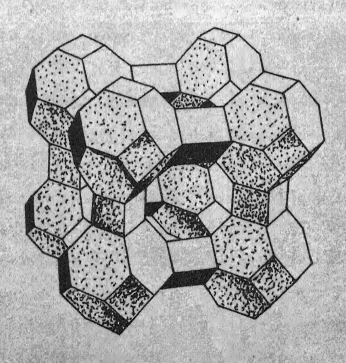


Fig. 2-4 - Arrangement of truncated cuboctahedron to give zeolite type A.

themselves enclose a second set of smaller cavities (β cages) o with an internal diameter of 6.6 A which are connected to the larger cavities by means of a distorted ring of six oxygen atoms of 2.2 A free diameter.

In each crystallographic unit cell of zeolite type A, there are 12 AlO, and 12 SiO, tetrahedra and therefore 12 monovalent cations. Eight of these sodium ions lie in the centre of the six rings in the hexagonal faces and four occupy positions adjacent to the eight membered ring. When the zeolite is hydrated, these four cations are completely hydrated and float in the centre of their coordination sphere of water molecules. When the water is removed, the cations are found to be located on the walls of the cavity [85]. The eight Na tions in the six rings are referred to as type I and the remaining four as type II. As discussed earlier, the zeolites readily exchange the alkali metal cations originally present for many other species of cations. the sodium ions in zeolite type A are replaced by calcium ions, the effective aperture size increases. The sodium form (effective aperture size 4.0 A) does not adsorb linear hydrocarbons but after one-third of the sodium ions are replaced by calcium (four per unit cell) many straight chain paraffins are readily adsorbed. This calcium form of zeolite A has an effective diameter of 5 A.

2.3 APPLICATIONS OF ZEOLITES AS ADSORBENTS:

One of the major industrial applications of zeolites is in the area of adsorption processes. Zeolite adsorbents are not only the most important adsorbents today, but their importance is increasing, mainly because of the following unique adsorptive properties:

- (a) Selective adsorption of molecules based on molecular dimensions;
- (b) Highly preferential adsorption of polar molecules;
- (c) Highly hydrophilic surface;
- (d) Variation of properties by ion exchange;
- (e) Large void volume and
- (f) Well defined crystalline structure.

Zeolite adsorbents are mostly used in a fixed bed. A number of columns packed with the zeolite adsorbent are interconnected with an automatic valve system to facilitate a continuous flow of industrial stream being processed. Each bed, however, goes through a stepwise cyclic operation, and during each cycle the adsorbed molecules in the zeolite bed are desorbed either by raising the bed temperature, lowering the bed pressure, displacing the adsorbate with another adsorbate, or a combination of these.

Important properties of zeolite adsorbents for a fixed bed application are adsorptive capacity and selectivity, adsorption—desorption rate, physical strength, low catalytic

activity, thermal-hydrothermal stability, chemical stability, and particle size and shape. The ultimate basis in selecting a zeolite adsorbent for a specific application would be the performance, price, and the projected service life of a product. Important industrial adsorption processes using zeolite adsorbents may be classified as follows:

- (i) hydrocarbon separation processes
- (ii) drying gases and liquids
- (iii) pollution control applications
 - (iv) separation and purification of industrial streams.

2.3.1 Hydrocarbon Separation:

2.3.1.1 n-Paraffin Separation:

n-paraffins need to be separated from gasoline to increase its octane number and from kerosene to use them as a raw material for biodegradable detergents. A detailed discussion on n-paraffin separation processes has been given by Ries [37].

Many workers [39, 42, 57, 59, 99] have reported that n-paraffins can be separated from gasoline to increase its octane rating, using molecular sieve 5A as an adsorbent. These processes mainly differ in the operating conditions during the adsorption cycle and the type of desorbent used to recover n-paraffins. Avery and Lee [5] separated n-paraffins $(C_5H_{12}-C_{12}H_{26})$ from a mixture of iso- and cyclic paraffins

present in naphtha and kerosene. Volf, et al. [100], could increase the octane number of gasolines of various boiling ranges by 17-20 points. Adsorption was carried out in the vapor phase at $25-30^{\circ}\mathrm{C}$ higher than the boiling point of the least volatile component in gasoline. Desorption was carried out by nitrogen at $300-350^{\circ}\mathrm{C}$. Galich, et al. [50] found that the adsorption of n-paraffins on Ca-A zeolite increased from C_7 to C_{10} paraffins then decreased. They reported that the time to reach equilibrium increased with molecular weight of alkane. Charles [27] proposed a countercurrent adsorption scheme and then desorption of n-paraffins by heating the loaded molecular sieve 5A at a temperature $320^{\circ}\mathrm{C}$.

Galich, et al. [51], have used steam as a desorbent. Investigations at ESSO [41] indicate that molecular sieve life is extended by using lower boiling dlefins rather than steam as a desorbent. With propylene as a desorbent, they did not observe any change in adsorption capacity after 16 cycles while with steam, the sieve capacity was reduced to two thirds of the fresh sieves after 3 cycles. Desorption with different hydrocarbons was tried by Fleck [45]. Mirskii, that adsorbed baraffins were desorbed et al. [73] suggested with an excess of a lower boiling n-paraffin. They preferred butane in place of pentane for displacing, adsorbed hexane and/or heptane because of its lower boiling point. James and Lewis [46] reported desorption by straight chain hydrocarbons with 1 to 3 fewer carbon atoms.

ESSO [43] has used ammonia as a desorbent and found that desorption efficiency increased with decrease in pressure. Carbon dioxide was also used by several workers [38, 48, 74] as a desorbent. Plachenov, et al. [84] reported that when carbon dioxide was used as desorbing agent the desorption of octane and hexane was 9-10 per cent higher than when nitrogen was used.

Afanas'ev, et al. [1] reported that the description of paraffins from zeolite NaX is more complete compared to that from CaA. Plachenov, et al. [85] compared the thermodisplacement and vacuum—thermal methods of description and found that former is more effective at temperatures less than 105° C while at higher temperatures both methods are equally effective. Areshidze and Chivadze [2] described the paraffins by increasing the temperature and reducing the pressure.

Kivitkovskii [63] reported that in case of n-heptanen-octane system larger amounts of n-octane (higher molecular weight paraffin) were adsorbed while n-nonane (lower molecular weight component) was preferentially adsorbed from n-nonanen-decane system.

Major commercial processes in n-paraffin separation are UOP's Molex process [21, 22, 23, 94], B.P's process [62, 105, 106], Exxon's Ensorb process [3, 86], Union Carbide's IsoSiv process [5, 54, 55], Texaco's T.S.F. process [31, 47], Shell's process [89], and VEB Leuna Werke's Parex process [104].

All processes, otherthan the Molex process, are in vapor phase and use fixed bed, cyclic technology. The processes are different, however, in operating pressure, temperature, desorption technique, and other operating conditions.

Desorption is generally carried out by displacement. Displacement agents mentioned in patented literature are generally low boiling n-paraffins, and ammonia or alkylamines. Steam a cheaper source has also been used as a desorbing agent.

Pressure swing operation is used for separating low carbon number n-paraffins. Purging, between adsorption and desorption cycles, is some times desirable when product purity is important.

In the Molex process [22, 23, 87] adsorption system consists of a single adsorption tower with multiple inlet-outlet points and a special rotary valve. The adsorption tower has many smaller adsorption chambers interconnected in series, and works under so called 'simulated moving bed' operation. Instead of moving the bed, the simulated moving bed operates by simultaneously advancing inlet-outlet points periodically.

2.3.1.2 Separation of p- and O-xylenes:

The rapid increase in the use of p-xylene as a raw material for polyester products necessitated the development of a process for separating p-xylenes from a mixture of C₈ aromatics. Broughton, et al. [24] and Atkins [4],in 1970, have developed a process for separating p-xylenes from a

mixture of C₈ aromatics containing xylenes and ethylbenzene. The process has been commercialized by the name U.O.P.'s

Parex process and the hardware for this process is similar

to Molex process for separation of n-paraffins. The process
is continuous liquid phase, simulated moving bed type. This

process can separate p-xylene from various types of feedstocks

with 99.5 per cent purity and recovery as high as 98.4 per cent

[24]. The high recovery is certainly an improvement over

the conventional crystallization processes. The adsorbent

used in U.O.P. process is a synthetic faujasite containing

cations of groups IA, IIA, or both [76-78]. U.S. Patents [14,
28] issued in 1972 also claim that sodium mordenite and

modified type-Y zeolite containing predominantly potassium

ions can separate p-xylenes from a mixture of p- and O-xylenes,
and a C₈ aromatic mixture, respectively.

2.3.1.3 Olefin Separation:

For the separation of olefins from a feedstock containing olefins and paraffins, the zeolite adsorbent used, according to patent literature, [79,80], is a synthetic faujasite with 1-40 weight per cent of atleast one cation selected from groups IA, IIA, IB, and IIB. The process is known as the U.O.P.'s Olex process. This process is also believed to use the same simulated moving bed operation in liquid phase as U.O.P.'s other hydrocarbon separation processes. Union Carbide's OlefinSiv process [7] is used to separate

n-butylenes from isobutylene. Product purities are claimed to be above 99 per cent for both n-butylene and isobutylene streams.

2.3.2 Drying Gases and Liquids:

Zeolites have a highly hydrophilic surface and are very efficient desiccants. Contrary to other non-zeolitic desiccants, such as silica gel and activated alumina, zeolite adsorbents have type I adsorption isotherm for water, which means a high water adsorption capacity at a low concentration. The zeolite type 3A, in particular, has the additional advantage of selective adsorption of water because of its small pore size. Although, in the past, alumina has been the most commonly used desiccant in drying cracked gas, 3A molecular sieve adsorbents have an overall economic advantage [82]. The main advantages of 3A molecular sieve over alumina and silica gel are its higher capacity and, therefore, smaller adsorption tower and its longer service time. Applications of zeolite adsorbents in drying other industrial gases and liquids have been discussed elsewhere [29, 30, 72].

2.3.3 Separation and Purification of Industrial Streams:

2.3.3.1 Purification of Air Prior to Liquefaction:

Liquefaction of air by cryogenic processes requires removal of water vapor and carbon dioxide to avoid heat exchanger freeze-up. A 13X molecular sieve has

been used for separation of water vapor and carbon dioxide from air in a single adsorption step. The 13X molecular sieves have not only higher adsorptive capacities but also faster rates of ${\rm CO_2}$ adsorption compared to type A. The optimum operating temperature for ${\rm CO_2}$ removal by 13X molecular sieve is reported as 160-190°K [102].

2.3.3.2 Natural Gas Purification:

Natural gas containing water vapor, carbon dioxide, hydrogen sulfide, etc. is purified by passing through either 4A or 5A molecular sieve bed. Other important applications of molecular sieves in natural gas purification [56, 58, 64,85] include purification of pipeline natural gas for liquefaction, drying natural gas prior to cryogenic hydrocarbon recovery using a turboexpander, and sweetening natural gas feed to ammonia plants.

2.3.3.3 Oxygen Enrichment of Air:

The increase in demand of oxygen or oxygen-rich air in biological waste water treatment plants generated a necessity for a low cost, on site oxygen generator. Other applications of oxygen enrichment of air are secondary smelting plants, river and pond aeration, pollution control in the pulp and paper industry, etc. Pressure swing adsorptive process using zeolite adsorbents [63, 66] is found to be advantageous over the conventional cryogenic air separation

process. Commercial processes known today are: Union Carbide process [33], the W.R. Grace process [63], the Bayer-Mahler process [13], the Nippon steel process [96, 95], and ESSO Research and Engineering processes [16, 44, 91, 92]. These processes differ in the type of zeolites used, operating pressures, number of adsorbent beds, and cyclic operating steps.

2.3.4 Pollution Control:

Zeolite adsorbents are effectively used to remove pollutants such as SO_2 , $\mathrm{H}_2\mathrm{S}$, and NO_X from industrial off-gas streams nearly at ambient temperature [52, 60, 65, 68]. Since water vapor usually exists along with these acidic compounds, an acid stable or acid resistant zeolite adsorbent is necessary for a long service life. Union Carbide offers three new processes for pollution control, namely, the PuraSiv-Hg process for mercury vapor removal, the PuraSiv-N process for NO_X removal from nitric acid plant off-gas, and the PuraSiv-S process for SO_2 removal from sulfuric acid plant off-gas.

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CHAPTER 3

EXPERIMENTAL RESULTS ON ADSORPTION OF PURE COMPONENTS

3.1 INTRODUCTION:

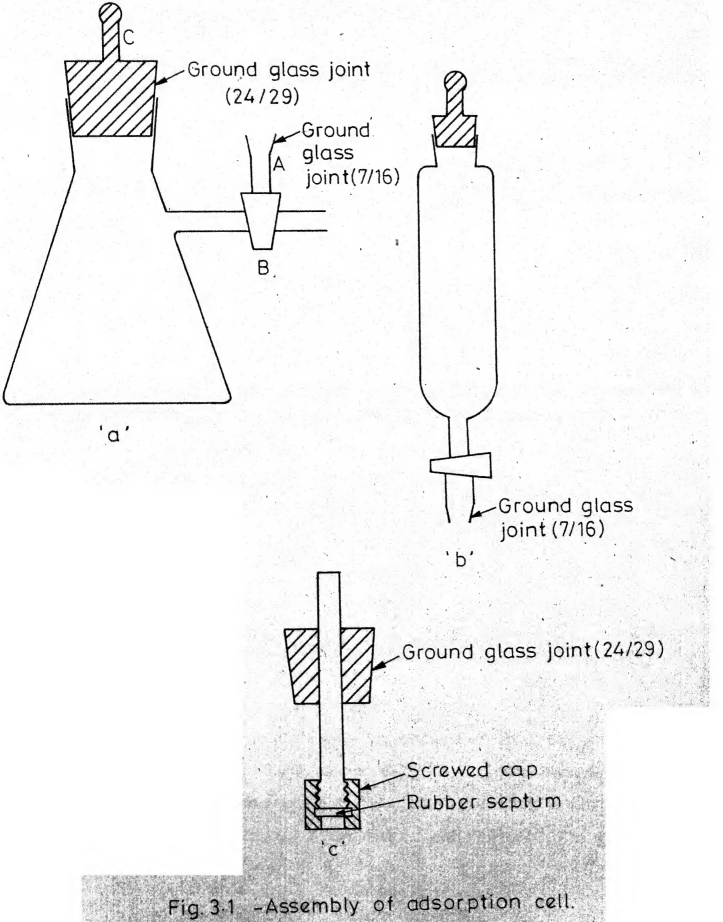
Demand for n-paraffins has been steadily increasing in pure form as well as mixtures because of a variety of novel applications. N=Octane is used in fermentation process for the manufacture of citric acid. Use of nparaffins in the manufacture of biodegradable detergents has increased very rapidly over past severalyears. sieve adsorption processes which are very effective, efficient and cheap are becoming popular for the separation of n-paraffins from petroleum fractions. For the design of adsorption columns, one needs the basic data on liquid phase adsorption of these n-paraffins on molecular sieves. Although several workers [4-8] have studied the vapor phase adsorption of pure and binary paraffins, practically no reliable data is available on liquid phase pure component adsorption of n-paraffins in published literature. Sundstrom and Krautz [9] have reported single component adsorption capacities for C7, C10, C12 and C14 n-paraffins. However, these authors have not given the adsorption isotherms, which are necessary for design. This has motivated undertaking the present study where adsorption isotherms have been measured for

 $^{\text{C}}_{5}$ to $^{\text{C}}_{8}$ n-paraffins using LMS-5A as an adsorbent. A temperature range of 6^{O} to 42^{O}C has been covered in this study.

3.2 EXPERIMENTAL:

The molecular sieves used were LMS-5A; 1/16'inch pellets containing 20 per cent inert clay binder. Benzene (A.R. grade, B.D.H.) and n-paraffins (G.R. grade; E. Merk/B.D.H.) had a minimum purity of 99.0 per cent. To study the effect of liquid concentration on the adsorption, pure n-paraffins were diluted with benzene. Due to its critical diameter (6.8 Å), benzene is essentially excluded from the pores (equilibrium loading of benzene on LMS-5A has been reported as 0.002 g/g zeolite [2]).

The experiments were conducted in a pyrex glass assembly, shown in Figure 3.1. which consisted of a 100 ml. adsorption cell attached to a 3-way vacuum stop cock. The molecular sieves were heated at 450°C for 48 hours and then transferred to the preheated cell maintained at 150°C which was then stoppered. The assembly containing the molecular sieves was then evacuated to 10⁻³ torr at 200°C for 4 hours to remove any water vapor or air left in the cell. The evacuated cell was then cooled to room temperature. A weighted quantity of liquid paraffin and benzene mixture was then added to the adsorption cell under vacuum with the help of the



assembly shown in Figure 3.1b which was connected to the cell at A (Figure 3.1a). Since the cell was under vacuum, the liquid sample was almost instantaneously sucked and the stopcock B (Figure 3.1a) was then closed. The adsorption cell was then transferred to a controlled temperature water bath with a shaker (Model G-86; New Brunswick Scientific Co., New Jersey) which was maintained at the desired temperature of adsorption within ± 0.25°C. Preliminary experiments were conducted with varying amounts of molecular sieves, shaker speeds and volume of sample. From these runs, nearly 30 g of molecular sieves, about 50 ml. of liquid sample and a shaker speed of 300 rpm were chosen as the operating conditions for the main runs.

The breakthrough curves were first determined to establish the time taken to attain equilibrium. To facilitate periodic withdrawal of liquid samples for chromatographic analysis, the assembly shown in Figure 3.1c was attached to the adsorption cell in place of stopper c (Figure 3.1a). This had a rubber septum through which the needle of a microlitre syringe could be pierced to reach the liquid in the cell. These samples were subsequently analysed on a thermal conductivity gas chromatograph (Chromatographic Instrumentation Company; Baroda). A 30 ft long column of 20 per cent bentone on chromosorb W was used for the analysis.

Detailed conditions for analysis are given in Appendix A3.1. Breakthrough runs were terminated when the composition of two successive samples were identical. In general, breakthrough point was reached in approximately 24 hours.

To ensure complete equilibration during adsorption studies, each run was continued for 48 hours and the liquid sample was then analysed to get the final concentration of n-paraffin in the solution. Each reported point on the adsorption isotherm was determined by averaging the results of two experiments conducted simultaneously in two identical cells.

3.3 RESULTS AND DISCUSSION:

3.3.1 Breakthrough Curves:

The effect of temperature on the breakthrough curve, for n-heptane is shown in Figure 3.2. Data are tabulated in Appendix A3.2). This figure shows that the time for reaching equilibrium decreases with increasing temperature. This is to be expected because adsorption is a mass transfer process and increasing the temperature increases the mass transfer coefficient, resulting in the breakthrough point being reached earlier.

3.3.2 Adsorption Isotherms:

Figures 3.3 to 3.6 (Data are tabulated in Appendices A3.3 to A3.6) show the adsorption isotherms for n-pentane,

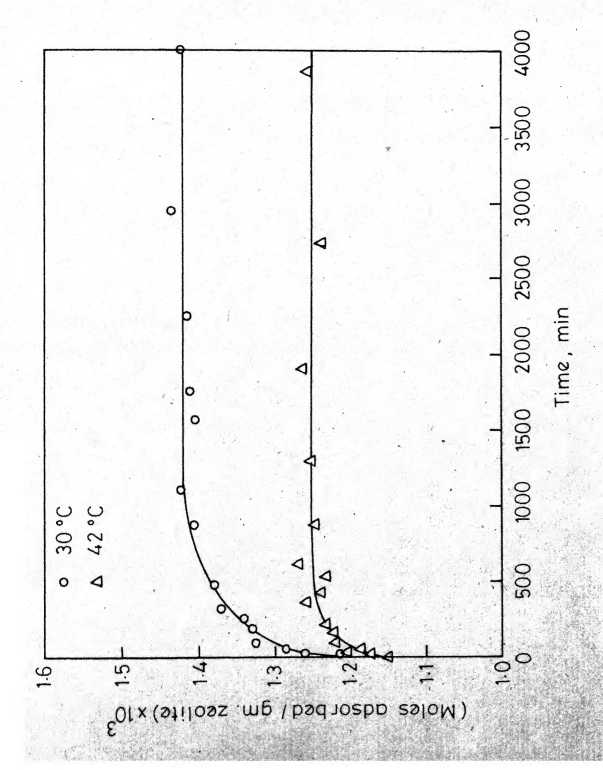


Fig. 3.2 - Breakthrough curves for n-heptane.

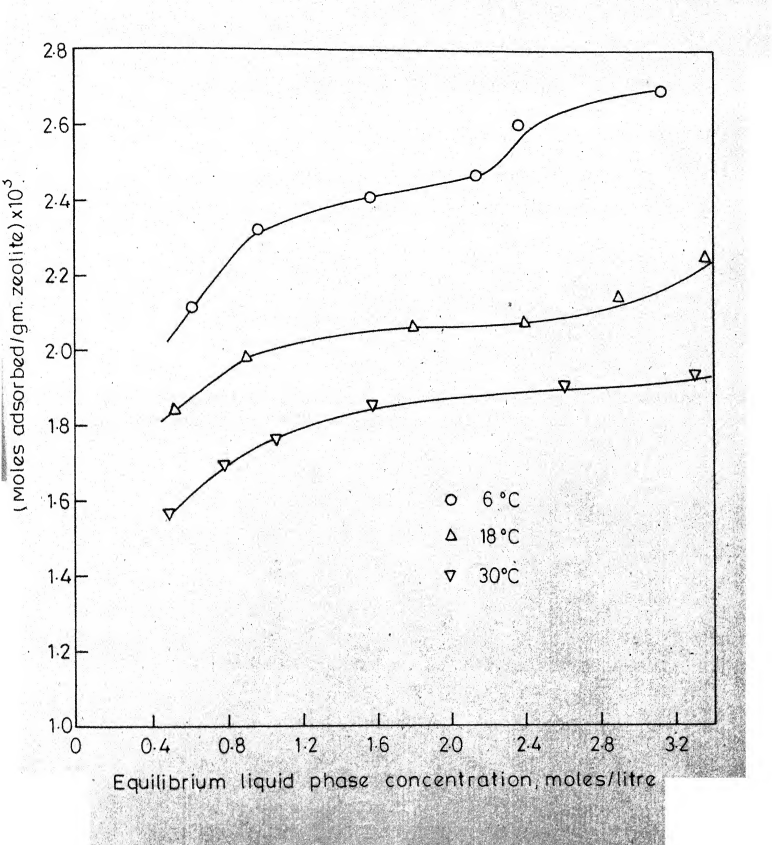


Fig. 3.3 - Adsorption isotherms of n-pentane.

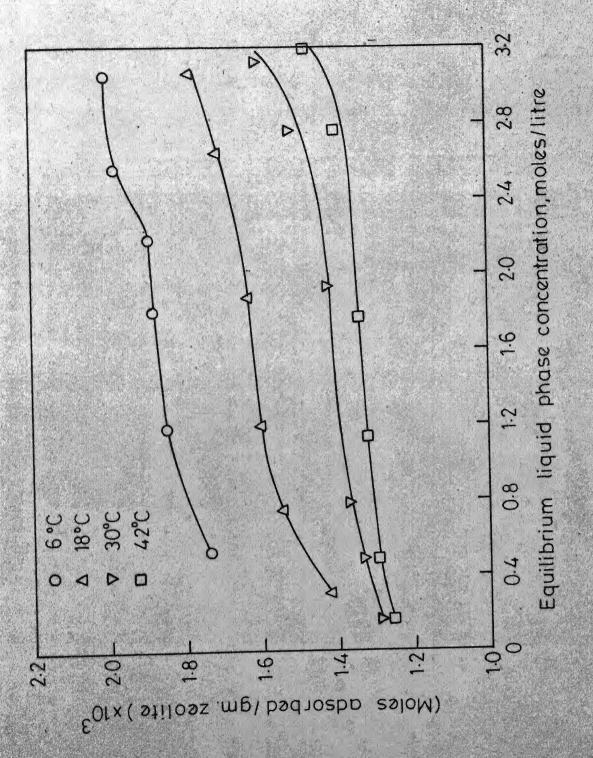


Fig. 3.4 - Adsorption isotherms of n-hexane.

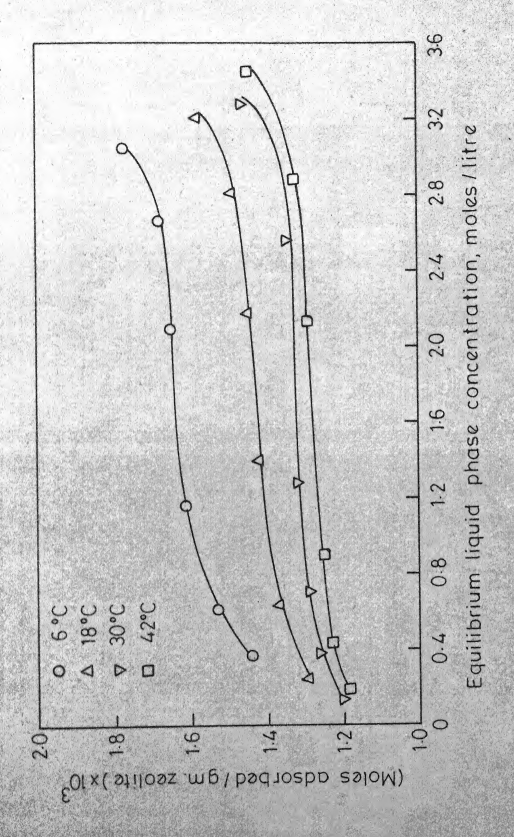


Fig. 3.5 - Adsorption isotherms of n-heptane.

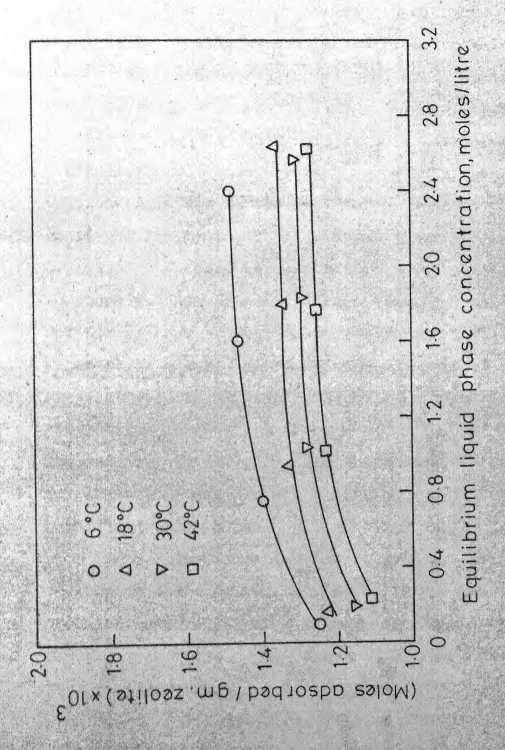


Fig. 3.6 - Adsorption isotherms of n-octane.

n-hexane, n-heptane and n-octane respectively. The general trend of the isotherms show an initial increase in equilibrium loading with concentration followed by a relatively large range of concentration over which the equilibrium loading is nearly constant and then a further increase at high concentrations. This upward trend at high paraffin concentration could not be observed for n-octane because of some practical limitations. This upward increase in equilibrium loading is probably associated with adsorption on external surfaces and not with intra-crystalline adsorption. Barrer and Lee [1] have also reported an upward inflexion in the vapor phase adsorption of hydrocarbons near the condensation temperature. As expected, at the same concentration, the equilibrium loading decreases with increasing temperature. A comparison of these Figures shows that the effect of temperature is maximum for n-pentane and reduces for higher paraffins, being minimum for n-octane in the present study.

For a particular temperature, the effect of concentration on the equilibrium loading is most significant for n-pentane and decreases with increase in molecular size.

A typical plot at 30°C is shown in Figure 3.7. This shows that as the equilibrium liquid paraffin concentration changes from 0.4 to 3.2 moles/litre, the equilibrium loading for n-pentane increases from 10.87 to 13.82 g/100 g zeolite which amounts to approximately 28 per cent. For n-heptane, the

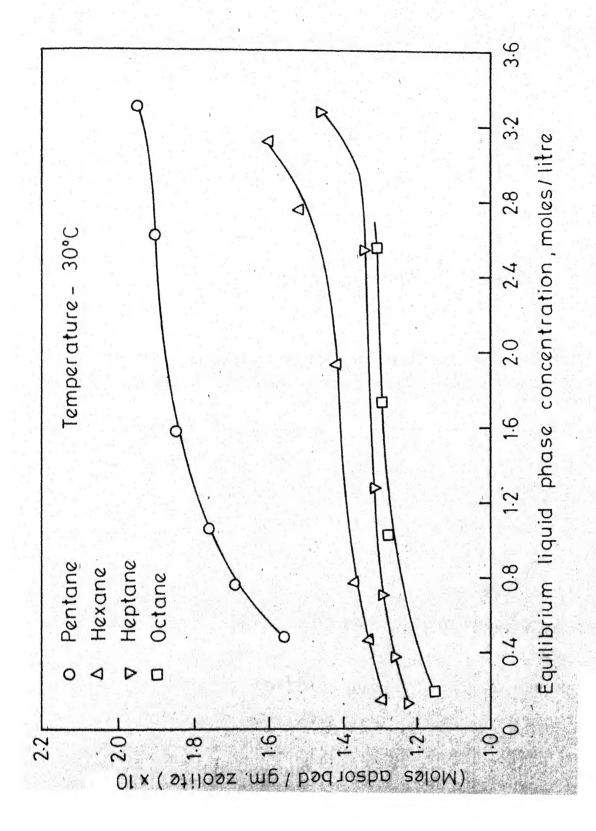


Fig. 3.7 -Adsorption isotherms of different n-paraffins at 30°C.

equilibrium loading increased from 12.50 to 14.30 g/loo g zeolite in the same range of concentration, which is only a 14.4 per cent increase. The results of equilibrium loading for n-heptane are in good agreement with those of Sundstrom and Krautz [9], who have reported a loading of 12.5 g/loo g zeolite at 30°C. These authors did not observe any change in the equilibrium loading with concentration; however, the concentration ranges have not been reported in their paper and it is possible that they did not cover as wide a concentration range.

For the same temperature and liquid concentration, the equilibrium loading decreases with increase in molecular size i.e. the lower molecular weight paraffin is preferentially adsorbed. With increasing molecular weight, the effect of size diminishes. Thus, at 18°C, the equilibrium loading for n-pentane is 2.03 x 10⁻³ moles/g zeolite at an equilibrium concentration of 1.6 moles/litre compared to 1.62 x 10⁻³, 1.43 x 10⁻³ and 1.34 x 10⁻³ moles/g zeolite for n-hexane, n-heptane and n-octane respectively. Peterson and Redlich [3] have also shown that liquid phase sorbability decreases with increasing number of carbon atoms for n-paraffins having 5 or more carbon atoms. This violates Traube's rule which states that higher molecular weight adsorbate should be preferentially adsorbed. The reason

for the preferential adsorption of lower molecular weight hydrocarbons on 5A molecular sieves is that in the zeolites both physico chemical adsorption and steric effects are important. Since type 5A molecular sieves have a pore of pening of 5A and n-paraffins have critical diameter of 4.9A, steric effect will play an important role. In shorter chain compounds (lower molecular weight n-paraffins) steric effects will be less in comparison to larger chain compounds.

In the present investigation, an experimental technique, for the liquid phase adsorption of hydrocarbons on zeolite has been perfected. Adsorption isotherms have been measured for different liquid paraffins in the temperature range 6 to 42°C. These data can be used to design single or multiple component adsorption plants.

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CHAPTER 4

EXPERIMENTAL RESULTS ON ADSORPTION OF MULTICOMPONENT SYSTEMS

4.1 INTRODUCTION:

n-Paraffins form an integral part of various petroleum fractions available in a refinery. In any fraction there are more than one component and any separation process has to be designed to remove mixture of these n-paraffins. molecular sieve process also different components will adsorb simultaneously in different proportions according to their relative affinity. In order to design these separation systems, multicomponent adsorption data is, therefore required. Very little information is available in published literature on multicomponent liquid phase adsorption of nparaffins on molecular sieves. Sundstrom and Krautz [3] have reported some data on the liquid phase adsorption of binary systems of C7, C10, C12 and C14 n-paraffins on LMS-5A. For all the systems studied, they found that the paraffin with the lower molecular weight was preferentially adsorbed and that the temperature had a negligible effect on the composition of adsorbate in equilibrium for a given liquid composition. Satterfield and Cheng [1] studied the adsorption of a wide variety of liquid binary hydrocarbon systems on NaY zeolite. Satterfield and Smeets [2] have reported the

liquid adsorption data on binary systems of n-octane either with n-decane, or n-dodecane, or n-tetradecane on NaY zeolite. They found that the lower molecular weight paraffin was preferentially adsorbed. They have suggested that the packing characteristics of the higher molecular weight paraffin, rather than the physico-chemical characteristic, play a dominant role in the adsorption selectivity on a zeolite.

In the present investigation, liquid phase adsorption of binary systems, namely, pentane-hexane, hexane-heptane, heptane-octane, pentane-heptane, and pentane-octane on synthetic zeolite LMS-5A have been studied. The ternary systems studied were pentane-hexane-heptane, pentane-hexaneoctane and pentane-heptane-octane. Benzene was used as a diluent in all the measurements since its adsorbability on the zeolite used is negligible. The equilibrium curves (mole fraction of lower molecular weight paraffin in liquid vs mole fraction of the same in the adsorbed phase) for all the binary systems were obtained by varying hydrocarbon concentration over a wide range. Measurements were made at 6°. 18°. 30° and 42°C to study the effect of temperature on the equilibrium curves. For all the binary systems, the effect of concentration and temperature on conventional separation factor, K, defined below was also investigated.

$$K = \frac{y_L/x_L}{y_H/x_H} \tag{4.1}$$

where \mathbf{x}_L and \mathbf{x}_H are molefractions of lighter and heavier paraffins in liquid phase and \mathbf{y}_L and \mathbf{y}_H are corresponding molefractions in adsorbed phase when the system is in equilibrium.

4.2 EXPERIMENTAL:

The experimental apparatus and procedure was the same as that for pure components described earlier (Chapter 3).

4.3 RESULTS AND DISCUSSION:

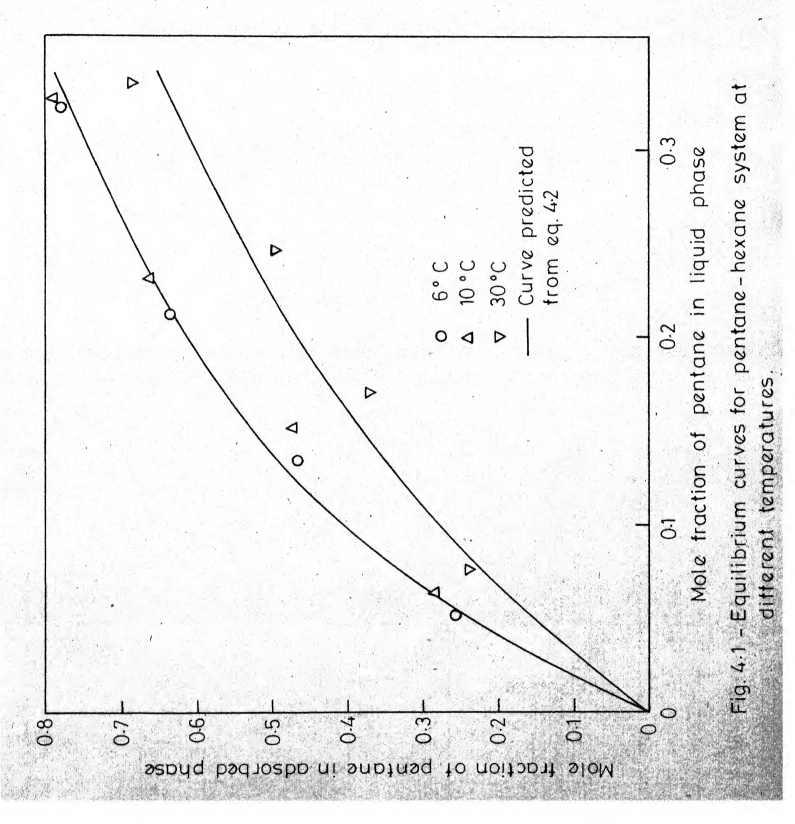
4.3.1 Equilibrium Curves for Binary Systems:

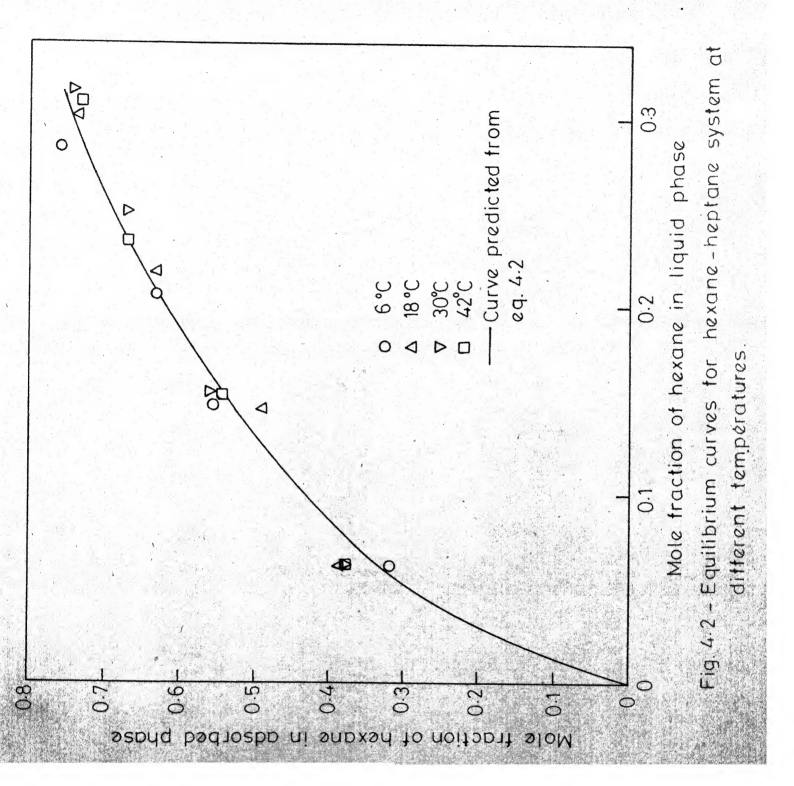
Figures 4.1 through 4.5 show the equilibrium curves for the various binary systems investigated (experimental data are tabulated in Appendices A4.1 through A4.4).

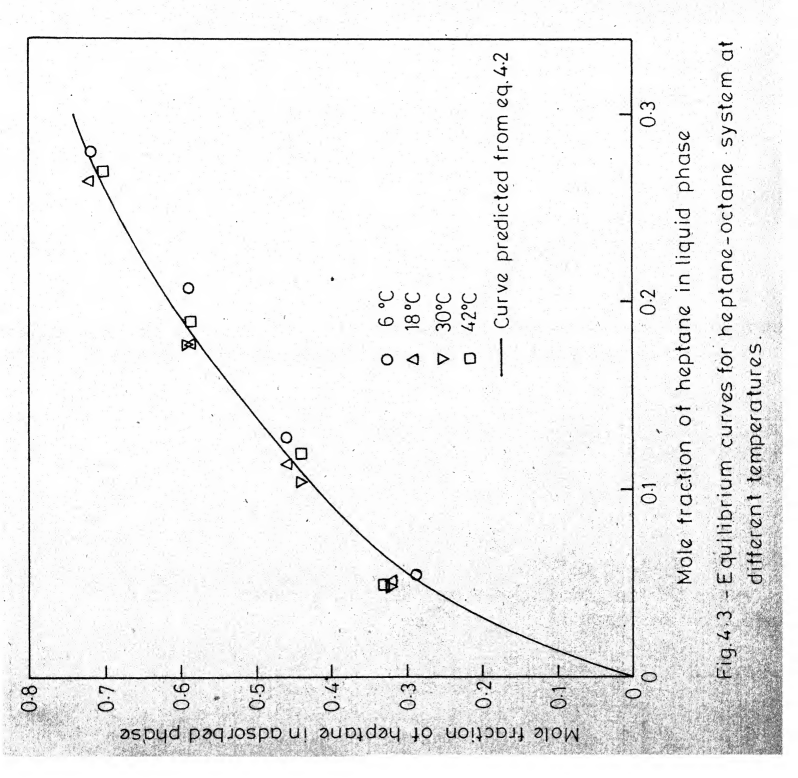
Temperature seems to have very little effect on equilibrium loading of hexane and heptane in hexane-heptane and heptane-octane systems as evidenced from Figures 4.2 and 4.3.

Sundstrom and Krautz [3] have also reported (for binary systems of higher paraffins) that temperature has a negligible effect on the composition of an adsorbate in equilibrium with a given liquid composition. There is a decrease in the amount of pentane adsorbed at 30°C compared to the lower temperatures for pentane-hexane, pentane-heptane and pentane octane systems (Figures 4.1, 4.4 and 4.5). Since T KANPUR

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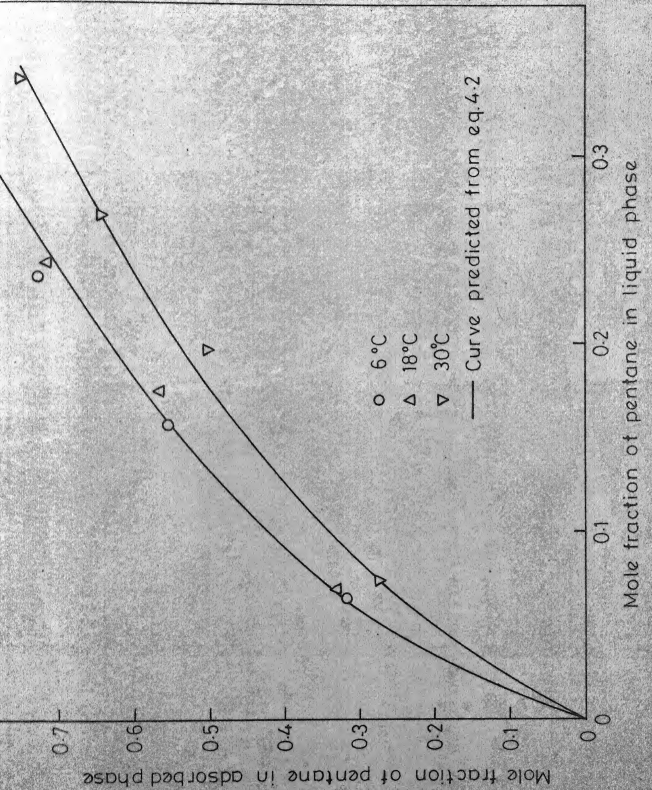
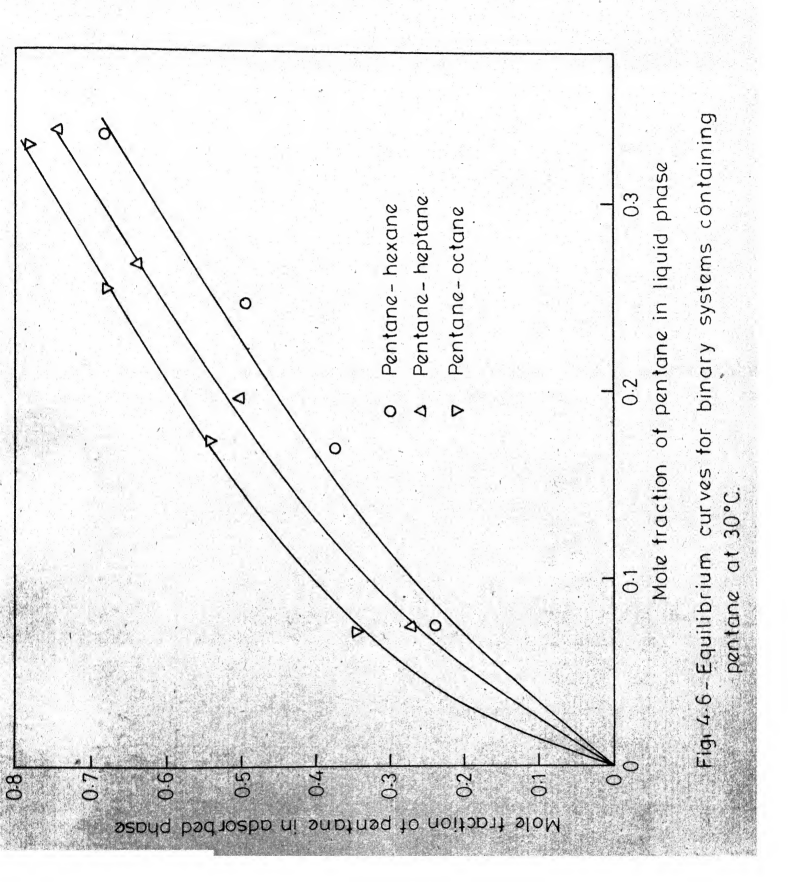


Fig. 4.4 - Equilibrium curvestor pentane-heptane system at

this temperature of 30°C is close to the boiling point of pentane, it is possible that due to the increased thermal motion, the probability of an adsorbed pentane molecule escaping from a cavity is higher, thus giving a lower adsorption of pentane at equilibrium. A comparison between Figures 4.1, 4.4 and 4.5 show that while for each system, equilibrium curves for 6° and 18° coincide, 30°C curve is distinct from 6° and 18° curve, separation increasing as we move from pentaneoctane system through pentane-heptane to pentane-hexane system. A possible explanation for this is that for pentanehexane system more unadsorbed space is left in the cavity in comparison to that in pentane-heptane system. resulting in more freedom of movement for the pentane molecule and hence more desorption of pentane occurs in pentane-hexane system. A plausible reason for negligible effect of temperature for pentane-octane system could be that with octane, pentane is so tightly packed leaving little or no space for free thermal motion and subsequent desorption. Equilibrium curves at 30°C. for binary systems containing pentane as one of the components are given in Figure 4.6. Adsorption of pentane is highest for pentane-octane system and is the least for pentane-hexane system. Satterfield and Smeets [2] have suggested that the heavier molecule finds its favourable packing while smaller molecule takes up the remaining space which it can do more readily because it is smaller. Similar



argument holds for the present case. Of the three systems shown in Figure 4.6, pentane-octane makes the most tight packing where as pentane-hexane system the least for the simple reason that there is maximum size difference in the former case and minimum in the later.

4.3.2 Separation Factors for Binary Systems:

Figures 4.7 to 4.11 show the variation of separation factor K, as a function of molefraction of lighter hydrocarbon in liquid phase on benzene free basis. However, the values of K will be the same even if the liquid phase mole fractions were defined inclusive of benzene. In all the cases, K increases with decrease in composition of lower molecular weight paraffin for all temperatures. Figure 4.7 shows that at 6° and 18°C, temperature has a negligible effect on K for pentane-hexane system. At 30°C, there is a drop in separation factor which may be because of desorption of pentane molecules at 30°C as already discussed earlier (sec. 4.3.1). Values of K were independent of temperature for all other systems except pentane-heptane where at 30°C the separation factor was lower than that at 6° or 18°C.

For all the systems studied, K was found to depend on the liquid phase concentration and assumed values less than or greater than one depending on this concentration. However, Sundstrom and Krautz [3] have reported K to be independent of

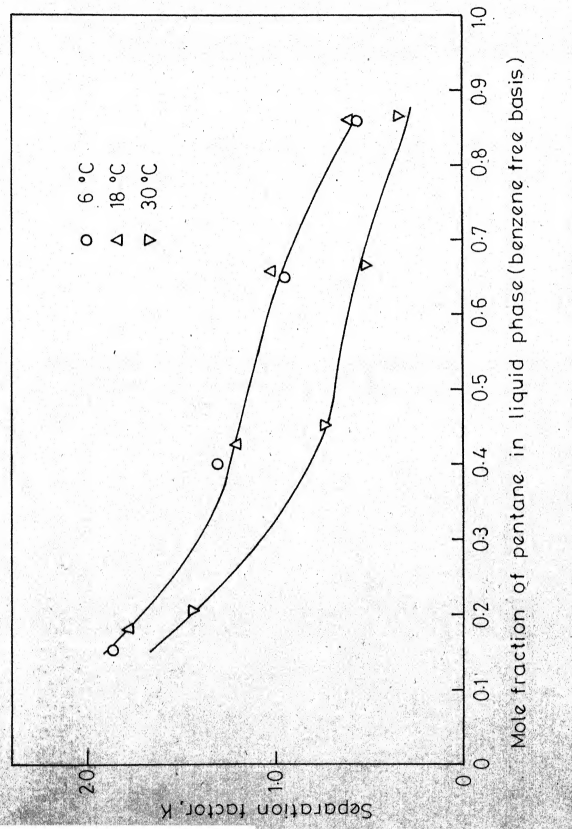
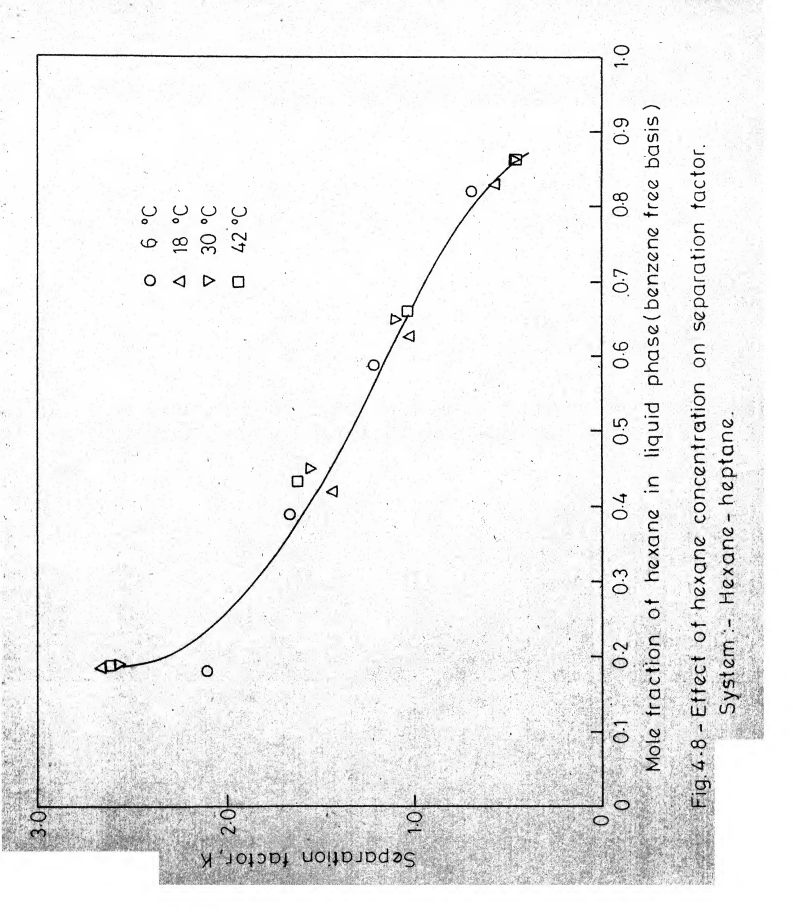
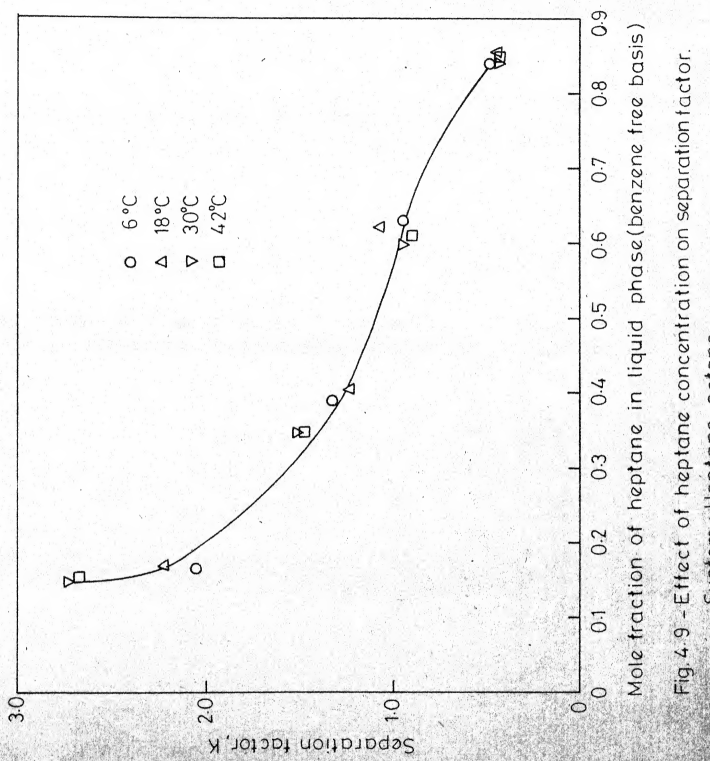
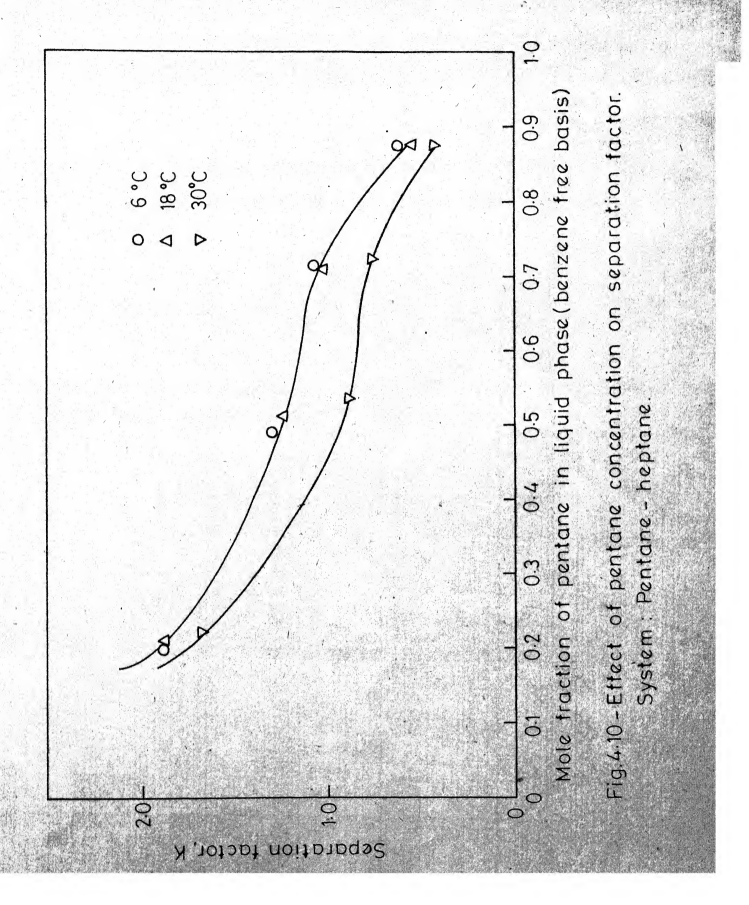


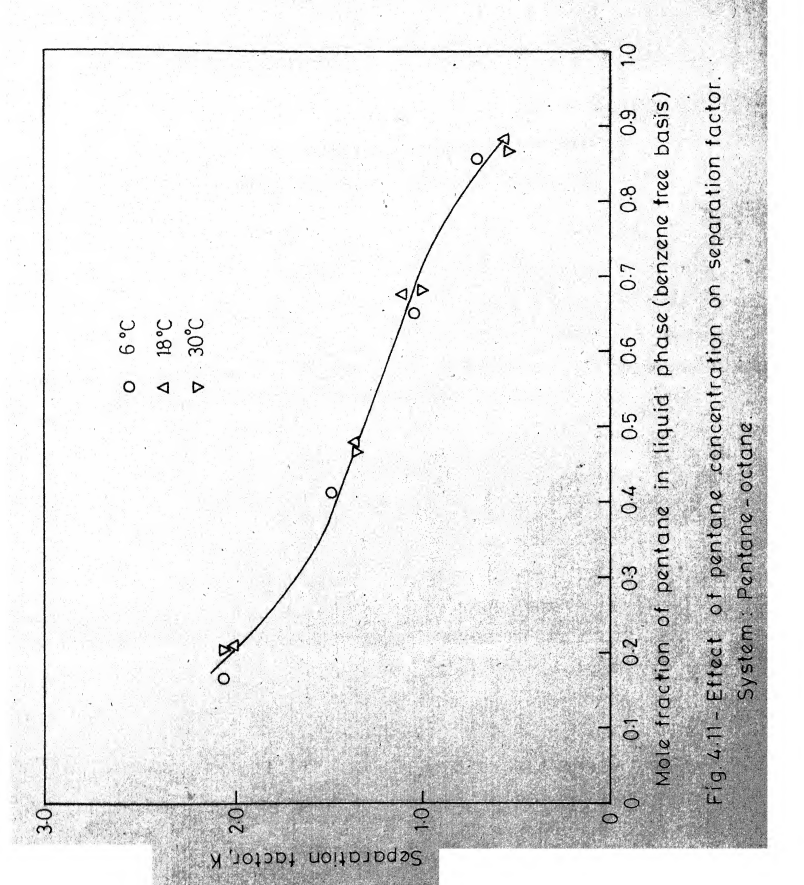
Fig. 4.7 - Effect of pentane concentration on separation factor. System :- Pentane-hexane.





System: Haptane-octane.





concentration and always greater than unity. The reason for this discrepancy becomes clear if one realises that in the present case (Figures 4.7 through 4.11) the abscissa does not represent the actual mole fraction of lighter paraffin as is normally understood. Usual practice is to define the mole fractions such that \mathbf{x}_L and \mathbf{x}_H will add to unity, whereas in this study, because of an inert component being present in the form of benzene, $\mathbf{x}_L + \mathbf{x}_H + \mathbf{x}_B = 1$; where \mathbf{x}_B is the mole fraction of benzene. However, by redefining the separation factor, it was possible to make it independent of concentration. The new separation factor \mathbf{K}_r was expressed as:

$$K_{r} = \frac{y_{L}/x_{L}}{y_{H}/(x_{H}+x_{B})}$$
 (4.2)

The values of K_{r} , for all the systems have been calculated and tabulated in Table 4.1. These values are the average of K_{r} calculated for different liquid phase concentrations at a fixed temperature. All values of K_{r} , thus calculated, are greater than unity. In Figures 4.1 through 4.5 the solid lines represent the calculated equilibrium curves using average values taken from this table in conjunction with eq. 4.2. Since the values of K_{r} at 30°C for the three binary systems that contains pentane as one component were much lower than those at other temperatures, the predicted curves have been drawn using values of

TABLE 4.1: SEPARATION FACTOR, K_r , FOR BINARY SYSTEMS

System		Separa	tion Factor	, K _r	
by s cent	6 ⁰ С	18°C	30°C	42 ⁰ C	Average
ВРХ	6.3	6.5	3.5ª	_	6.4
BXH	6.8	6.9	6.5	6.6	6.7
ВНО	6.8	7.3	6.6	6.8	6.9
ВРН	7.2	7.4	5.0 ^a	, ,	7.3
BPO	7.6	7.4	6.8 ^a	<u>-</u>	7.5

^aThose values have not been included in calculating the average values of ${\tt K}_{\tt r}{\mbox{.}}$

Notes: BPX = Pentane-heXane

BXH = heXane-Heptane

BHO = Heptane-Octane

BPH = Pentane-Heptane

BPO = Pentane-Octane

 $K_{\mathbf{r}}$ at this temperature rather than average values. The fact that these predicted curves represent the experimental data well is the proof that the redefined $K_{\mathbf{r}}$ is indeed independent of concentration. For binary systems containing pentane, the average $K_{\mathbf{r}}$ values increased with an increasing molecular weight of the heavier component. This shows that separation is easiest, for the systems studied, in pentane-octane system and most difficult in pentane-hexane system.

4.3.3 Adsorption in Ternary Systems:

The three ternary systems studied were pentane-hexane-heptane, pentane-hexane-octane and pentane-heptane-octane at 6°, 18° and 30°C. Adsorption results are presented in Tables 4.2 through 4.4 (operating conditions are tabulated in Appendices A4.5 to A4.7). Total number of molecules adsorbed per cavity decreased with increase in temperature. It was not possible to define a separation factor as was done for binary systems and these data were used to validate the multicomponent adsorption model as discussed later (Chapter 5).

SYSTEMS AT 60C TABLE 4.2: EXPERIMENTAL RESULTS FOR TERNARY

Run	Equilibrin	Equilibrium composition liquid phase, mole frac	tion in fraction	Molecules ausoi bou cavi ty	cavi ty	Tod no	molecules adsorbed
	1	2	3	1	2	3	per caving
PXH-1	0,235	0.047	0.083	3.29	1.26	1.29	5.84
PXH-2	0.082	0.174	0.089	1.45	2,46	1.30	5.21
PXH-3	0.053	0.139	0.165	1.20	2.02	1.72	4.94
PXH-4	0.134	0.087	0.141	1.99	1.61	1.60	5.20
PX0-1	0.207	0.046	0.045	3.49	1.27	1,15	5.91
PX0-2	0.082	0.149	0.045	1.67	2.50	1.13	5.30
PX0-3	0.052	0.132	980.0	1.35	2,26	1.47	5,08
PX0-4	0.124	0.088	0.088	2,11	1.64	1.44	5.19
PH0-1	0.223	0.056	0.050	3.60	1,16	1.19	5.95
PH0-2	0.082	980.0	980.0	1,66	1.44	1.62	4.72
PH0-3	0.051	0.162	0.045	1.32	2.17	1.23	4.72
PH0-4	0.055	0.130	0.075	1.35	1.83	1,51	4.69

the heaviest hydrocarbon is the lightest hydrocarbon in given ternary system and 3 is Н

						The second secon	THE RESIDENCE AND ADDRESS OF THE PARTY OF TH
Run	Equilibrium liquid phase	1 (1)	Composition in , mole fraction	Molecul	Molecules adsorbed cavity	ed per	Total number of molecules adsorbed
	-	2	3	1	2	3	per caviny
PXH-5	0.113	0.051	0.165	1,66	1.17	1.86	4.69
РХН-6	0.119	0.123	0.108	1.62	1,61	1.37	4.60
PXH-7	0,216	0.121	0.028	2.74	1.54	66*0	5.27
РХН-8	0.129	0.168	0.029	1,85	2.04	86 0	4.87
PX0-5	0.045	0.048	0.170	1.22	1,23	1.96	4.41
PX0-6	0.221	0.073	0.023	3.15	1.31	1.03	4.49
PX0-7	0.114	0.144	0.040	1,80	2.04	1.07	4.91
PX0-8	0.056	0.153	0.089	1,21	2.07	1.29	4.57
PHO-5	0.101	0.042	0.143	1.70	1.09	1.73	4.52
PH0-6	0.216	0.044	0.025	3.42	1.11	1.02	5.55
PHO-7	0.063	0,103	0.092	1.33	1.58	1.38	4.29
PH0-8	191.0	660*0	0.036	2,47	1.46	1.06	4.99
		the statement of the st	STOP PROBLEM PROBLEM PROBLEM STORES				
			1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1				6

Note: Notations are same as in Table 4.2

TABLE 4.4: EXPERIMENTAL RESULTS FOR TERNAHY

Run	Equilibrio 11, quid	Equilibrium composition in liquid phase, mole fraction	tion in	Molecul ce	Molecules adsorbed per cavity	bed per	Total number of molecules adsorbed
		2	3	1	2	3	per caving
PXH-9	0.077	0.221	0.044	1.04	2.54	1.04	4.62
PXH-10	0.260	0.055	0.042	2.28	1.32	1.12	4.72
PXH-11	0.184	0.135	0.046	1.54	1.90	1.08	4.52
PXH-12	0.170	0.056	0.103	1.67	1.30	1.42	4.39
PXH-13	0.072	0.144	0,115	1.09	1.96	1,30	4.35
PXH-14	0.123	0.037	0.181	1.43	1.16	1.80	4.39
PXH-15	0.142	0.111	0.083	1.44	1.78	1.24	4.46
PXH-16	0.150	0.138	0.103	1.38	1.80	1,23	4•41
1	1 1 1 1 1			1 1 1	1 1 1 1	I I I I	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
PX0-9	0.071	0.216	0.050	1.06	2.62	66*0	4.67
PX0-10	0.165	0,113	0.045	1.62	1.96	76.0	4.55
PX0-11	0.254	0.041	0.049	2.59	1.30	1.02	4.91
PX0-12	0.168	0.044	660.0	1.90	1.36	1.25	4.51

Table 4.4 (contd)

Run	Equilib Liquid	Equilibrium composition in Liquid phase, mole fraction	Equilibrium composition in Liquid phase, mole fraction	Molecu	Molecules adsorbed per cavity	bed per	Total number of molecules adsorbed
	7	2	3	7	2	3	per cavity
PH0-9	0.070	0.043	0.185	1,30	1,16	1,87	4.33
PH0-10	0.160	0.101	0.055	1.91	1.56	1,09	4.56
PH0-11	0.168	0.046	0.114	2.01	1.18	1,30	4.49
PHO-12	0.270	0.039	0.054	2.84	1.08	1.05	4.97

Note: Notations are same as in Table 4.2

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CHAPTER 5

MATHEMATICAL MODEL FOR ADSORPTION OF PURE AND MULTICOMPONENT HYDROCARBON SYSTEMS

5.1 INTRODUCTION:

The phenomenon of adsorption has been widely studied and extensive experimental data are now available for pure component vapor phase adsorption of light hydrocarbons and other compounds on different zeolites. However, industrial separation processes generally involve multicomponent adsorption, data for which are scanty. In any petroleum fraction, there are large number of combinations of hydrocarbons and it is, therefore, not possible to experimentally study all these systems. Hence, for the design of any adsorption process, a reliable method for predicting equilibrium adsorption data for mixtures from the isotherm of the pure components is essential.

The conventional adsorption models (Langmuir, Volmer, etc.) are not applicable to zeolite adsorption because the basic assumptions from which the models are derived are generally not fulfilled [2,3,19]. Other available approaches are potential theory and thermodynamic methods. According to the Dubinin-Polanyi potential theory [20], potential curves for adsorption of different gases, measured at different values of temperature and surface coverage on the same adsorbent, may be coalesced into a single curve

by a suitable choice of the coefficient of affinity. The usual choice for this coefficient is the molar volume of the liquified adsorbate. This property of pure-gas adsorption has been used by various investigators [6, 11,12] to predict adsorption equilibria for gas mixtures. The key assumption is that the characteristic potentials for various components of the mixture, and hence their fugacities in the gas phase are given by characteristic curves for adsorption of the pure gases. The extensions of the Dubinin-Polanyi theory to the case of mixtures is valid only when the surface potentials of pure adsorbates are equal at the state of saturated vapor [20]. The thermodynamic methods are those that are independent of any particular theory of the adsorption. Thermodynamic methods of estimating adsorption equilibria of gas mixtures include those of Myers and Prausnitz [13], Kidnay and Myers [8], Lewis, et al. [11], Arnold [1] and Cook and Basmadjian [5]. The basic postulate, common to all these models, is that the adsorbed phase forms an ideal adsorbed solution, but the methods differ in their choice of standard states. However, only the method of Myers and Prausnitz [13] is thermodynamically consistent and hence extensively used.

The structural regularity of zeolite lattices and the existence of well defined cavities within which the sorbate molecules are occluded, suggests that it should be possible

to develop a satisfactory theoretical equilibrium isotherm from the principles of statistical thermodynamics. Ruthven and co-workers, using this approach have proposed pure component [14] and binary gas phase [15-18] models. Binary model proposed by them is valid only for low concentrations.

No attempt has been made so far to develop theoretical models for adsorption in liquid phase even for pure components. However, since in the present study, liquid phase adsorption of single and multicomponent hydrocarbon systems has been investigated, for reasons already mentioned, it is necessary that a liquid phase prediction procedure is developed. no clues are available in literature on starting with liquid phase adsorption models, in this first attempt, use has been made of vapor-phase theory. It has been assumed that adsorption from liquid phase will be the same as that from vapor phase if this vapor is in equilibrium with the liquid. In otherwords, if we take a closed vessel partially filled with the liquid adsorbate, a vapor phase will be formed which will reach equilibrium with the liquid after sufficient time has elapsed. If an adsorbent is now placed in this closed system either in liquid phase or in vapor phase, the adsorption will be the same. It is assumed that if the adsorbent is placed in vapor phase, adsorption will disturb the equilibrium temporarily but molecules will come out of the liquid phase and restore vapor phase equilibrium almost

instantaneously. This assumption is usually valid since adsorption at vapor-liquid interface is negligible. This implies that one can use a vapor phase adsorption model to predict liquid phase adsorption if use is made of saturation vapor pressure at the given temperature in place of pressure for pure components. In case of multicomponent systems, partial pressures are obtained using Raoult's law since mixture of paraffins can be taken as an ideal solution.

5.2 MATHEMATICAL MODEL FOR ADSORPTION OF PURE n-PARAFFINS:

Consider a system of zeolite lattice with M cavities and an adsorbate. Because of the structural regularity of the zeolite it may be assumed that these cavities are separate subsystems each of which can adsorb any number of molecules s, from zero to a maximum of m. The canonical ensemble partition function for the system is [7]

$$Q = \sum_{s=0}^{m} q(s) \lambda^{s}, \qquad \lambda = e^{\mu/kT}$$
 (5.1)

where q(s) is the partition function for a cavity containing s sorbate molecules, μ is the chemical potential, k is the Boltzmann constant and T is the absolute temperature. When all the sites are equivalent (as in molecular sieves), the macroscopic system of M cavities may be regarded as a grand ensemble of subsystems. The grand partition function for this case is

$$G = Q^{M} = \begin{bmatrix} \sum_{s=0}^{m} q(s) & \lambda^{s} \end{bmatrix}^{M}$$
 (5.2)

The average number of molecules for this macroscopic system is given by [7]

or
$$C = \frac{\overline{N}}{M} = \lambda \frac{\partial \ln G}{\partial \lambda} = M \lambda \frac{\partial \ln Q}{\partial \lambda}$$
$$C = \frac{\overline{N}}{M} = \lambda \frac{\partial \ln Q}{\partial \lambda} = \frac{\sum_{s=0}^{m} q(s) \lambda^{s}}{\sum_{s=0}^{m} q(s) \lambda^{s}}$$

where C is the average number of molecules adsorbed per cavity.

The above equation can be written in terms of configuration integrals Z(s), as

$$C = \frac{\sum_{s=0}^{m} s Z(s) a^{s}}{\sum_{s=0}^{m} Z(s) a^{s}}$$
(5.3)

where Z(s) is the configuration integral for a subsystem of s-sorbate molecules occluded within a single zeolite cavity and is defined as

$$Z(s) = \frac{1}{s!} \int_{V} \exp\{-U_s(r_1...r_s)/kT\} dr_1...dr_s$$
 (5.4)

where r_i is the position vector, v is the volume of the zeolite cavity, U_s is the potential energy for the subsystem, a is the activity of the sorbate (= p/kT) for an ideal gas.

When there is no sorbate molecule occluded, we get

$$Z(0) = 1$$

and for one molecule,

$$Z(1)$$
 a = αp

where α is the Henry's constant and p is the pressure. Expansion of equation (5.3) gives

$$C = \frac{Z(1)a + 2Z(2) a^{2} + \dots + m Z(m) a^{m}}{1 + Z(1)a + Z(2) a^{2} + \dots + Z(m) a^{m}}$$
(5.5)

Equation (5.5) defines the adsorption isotherm for the system in terms of configuration integrals defined by equation (5.4). In order to compare theory with experiment, it is necessary to express configuration integrals in terms of measurable quantities. The following approximations are made.

- (i) It is assumed that the form of the potential field within a zeolite cavity is independent of the number of molecules occluded within the cavity.
- (ii) When two or more molecules are present within a given cavity, it is assumed that the molecules move randomly and independently in the potential field resulting from the combined effects of sorbate-sorbent and sorbate-sorbate potentials.

- (iii) The effect of sorbate molecules in neighbouring cavities is neglected.
- (iv) The energy of interaction between sorbate molecules in the same cavity is represented by Sutherland potential, defined as

$$U_{s}(r) = \infty$$
, $r < \sigma$; $U_{s}(r) = -\epsilon(\sigma/r)^{6}$, $r \ge \sigma$

where ε and σ are the molecular constants. With these approximations the configuration integrals may be written as

$$Z(s) = \frac{Z(1)^{s}}{s!} \left(1 - \frac{s \beta}{v}\right)^{s} \exp\left(\frac{s \beta \epsilon}{v k T}\right), 2 \le s \le v/\beta$$
(5.6)

where $\beta = \frac{2}{3} \pi \sigma^3$ is the effective co-volume of the sorbate. In this expression, the factor (1-s β /v) accounts for the reduction in free volume of the cavity while the exponential term represents the increase in adsorption potential due to the mutual attraction of the sorbate molecules. Substitution of eq.(5.6) in eq. (5.5) gives

$$Z(1)a + Z(1)^{2} a^{2} (1-2 \beta/v)^{2} \exp \left(\frac{2 \beta \epsilon}{v k T}\right) + \dots +$$

$$C = \frac{Z(1)^{S} a^{S}}{(m-1)!} \left(1 - \frac{m \beta}{v}\right)^{m} \exp \left(\frac{m \beta \epsilon}{v k T}\right)$$

$$1+Z(1)a + \frac{Z(1)^{2}a^{2}}{2!} (1-2\beta/v)^{2} \exp \left(\frac{2 \beta \epsilon}{v k T}\right) + \dots +$$

$$\frac{Z(1)^{S} a^{S}}{m!} \left(1 - \frac{m\beta}{v}\right)^{m} \exp \left(\frac{m \beta \epsilon}{v k T}\right)$$

Substitution of $Z(1)a = \alpha p$ in above equation gives

$$\alpha p + (\alpha p)^{2} (1-2\beta/v)^{2} \exp \left(\frac{2 \beta \epsilon}{v k T}\right) + \dots + \frac{(\alpha p)^{m}}{(m-1)!} (1-m \beta/v)^{m} \exp \left(\frac{m \beta \epsilon}{v k T}\right)$$

$$1+\alpha p + \frac{(\alpha p)^{2}}{2!} (1-2 \beta/v)^{2} \exp \left(\frac{2 \beta \epsilon}{v k T}\right) + \dots + \frac{(\alpha p)^{m}}{m!} (1-m \beta/v)^{m} \exp \left(\frac{m \beta \epsilon}{v k T}\right)$$
(5.7)

This is the theoretical equation for the isotherm for a pure component in terms of Henry's constant, α , and effective molecular volume, β. Some difficulty arises in the choice of appropriate value for eta, since the molecular volume of the sorbate, and hence the saturation capacity of the zeolite, varies with temperature. Dubinin [16] has suggested that the molar volume of the intracrystalline fluid may be approximated by a linear interpolation between the molar volume of the saturated liquid sorbate at its normal boiling point, TR, and the vander Waals co-volume at the critical temperature, T_C. This approximation has been used by Ruthven and co-workers [14-18] who found it valid only at very low adsorption. With this approximation eq. (5.7) is left with a single parameter α which, in principle, may be calculated, in terms of molecular constants, by evaluating the configuration integrals for a single sorbate molecule

occluded within a cavity. However, such calculations are only approximate [17, 18] since configuration integrals cannot be exactly calculated. In practice, α can either be calculated by the slope of the isotherm at low concentration or by matching the theoretical equation to an experimental isotherm. In general, it has been found [16] that the exponential term in eq. (5.7) has a very minor effect. This factor can, therefore, be omitted to give a simpler equation for the isotherm as

$$C = \frac{\alpha p + (\alpha p)^{2} (1-2 \beta/v)^{2} + \dots + (\alpha p)^{m} (1-m\beta/v)^{m}/(m-1)!}{1 + \alpha p + \frac{(\alpha p)^{2}}{2!} (1-2 \beta/v)^{2} + \dots + \frac{(\alpha p)^{m}}{m!} (1-m \beta/v)^{m}}$$
(5.8)

This expression is much simpler in comparison to eq.(5.7) and can be handled without much difficulty. Since benzene was used as a diluent, even in pure component systems there were two components in the liquid phase. Application of Raoult's law gives

$$C = \frac{\alpha Px + (\alpha Px)^{2} (1-2\beta/v)^{2} + ... + \frac{(\alpha Px)^{m}}{(m-1)!} (1-m \beta/v)^{m}}{1 + \alpha Px + \frac{(\alpha Px)^{2}}{2!} (1-2 \beta/v)^{2} + ... + \frac{(\alpha Px)^{m}}{m!} (1-m \beta/v)^{m}}$$
(5.9)

where P is the vapor pressure of the paraffin at adsorption temperature and x is the equilibrium mole fraction of the same in liquid phase.

Equation (5.9) along with the constraint m $\beta \leq v$ was used with pure component adsorption data to give values of α and β .

5.3 EXTENSION OF THE MODEL TO BINARY SYSTEMS:

The pure component model developed above has been extended for binary systems. In binary systems, there are, two adsorbates and both of them can be occluded individually or simultaneously in any cavity of the zeolite. There will be competition between the two components and the one which has a higher affinity for the adsorbent will be adsorbed preferentially in comparison to the other. The total number of molecules of both components will be restricted by the volume of the cavity. For a binary mixture of components A and B, the grand partition function, G_{AB} [17] is

$$G_{AB} = \begin{bmatrix} \sum_{j=0}^{\infty} & \sum_{i=0}^{\infty} & q(i,j) & \lambda_A^i & \lambda_B^j \end{bmatrix}^{M} = Q_{AB}^{M}$$
 (5.10)

where Q_{AB} is the canonical ensemble partition function for the system containing components A and B, q(i,j) is the partition function for a cavity containing implecules of A and j molecules of B, $\lambda_A = e^{\mu_A/kT}$ and $\lambda_B = e^{\mu_B/kT}$ (μ_A and μ_B are the chemical potentials of components A and B and T is the absolute temperature). The summation in the above equation is carried out overall values of i and j given by

$$i \beta_A + j \beta_B \leq v$$
 (5.11)

where β_A and β_B are the effective molecular volumes of A and B respectively at adsorption temperature. In terms of configuration integrals, eq.(5.10) can be written as

$$G_{AB} = \begin{bmatrix} \mathbf{\hat{\Sigma}} & \mathbf{\hat{\Sigma}} & \mathbf{\hat{Z}(i,j)} & \mathbf{\hat{a}_{A}} & \mathbf{\hat{a}_{B}}^{\mathbf{j}} \end{bmatrix}^{\mathbf{M}} = \mathbf{Q}_{AB}^{\mathbf{M}}$$
 (5.12)

where Z(i,j) is the configuration integral for system containing i molecules of A and j molecules of B and a_A and a_B are the activities of component A and B respectively. Average number of molecules of A and B adsorbed per cavity (C_A and C_B) are given by

$$C_{A} = \frac{n_{A}}{M} = \lambda_{A} \left(\frac{\partial \ln Q_{AB}}{\partial \lambda_{A}} \right)_{\lambda_{B}, T}$$
 (5.13)

$$C_B = \frac{n_B}{M} = \lambda_B (\frac{\partial \ln Q_{AB}}{\partial \lambda_B})_{\lambda_A} T$$
 (5.14)

Substitution of eq. (5.12) in eqs. (5.13) and (5.14) gives

$$C_{A} = \frac{\sum_{j=1}^{\sum_{i}} i Z(i,j) a_{A}^{i} a_{B}^{j}}{\sum_{j=1}^{\sum_{i}} Z(i,j) a_{A}^{i} a_{B}^{j}}$$
(5.15)

$$C_{B} = \frac{\sum_{j=1}^{N} j Z(i,j) a_{A}^{i} a_{B}^{j}}{\sum_{j=1}^{N} \sum_{j=1}^{N} Z(i,j) a_{A}^{i} a_{B}^{j}}$$
(5.16)

The configuration integrals for two components, with the same

assumptions as given for pure components (sec.5.2) are given by

$$Z(i,j) = \frac{Z(1,0)^{\frac{1}{2}} Z(0,1)^{\frac{1}{2}}}{i! j!} \left(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v}\right)^{\frac{1}{2}+\frac{1}{2}} \exp\left(\frac{i\beta_{A} \varepsilon_{A} + j\beta_{B} \varepsilon_{B}}{v k T}\right)$$

$$(5.17)$$

(where ε_A and ε_B are molecular constants for components A and B respectively) with the condition of eq.(5.11). Z(1,0) is the configuration integral when only one molecule of A and no molecules of B is adsorbed i.e.

$$Z(1,0) = Z_{\Lambda}$$
 (5.18)

Similarly,

$$Z(0,1) = Z_{R}$$
 (5.19)

$$Z(0,0) = Z(0) = 1$$
 (5.20)

Therefore.

$$Z(1,0) a_{A} = Z_{A} a_{A} = \alpha_{A} p_{A}$$
 (5.21)

and

$$Z(0,1) a_B = Z_B a_B = \alpha_B p_B$$
 (5.22)

where α_A and α_B are Henry's constants for components A and B respectively and p_A and p_B are partial pressures of A and B respectively. Expansion of eq.(5.15) gives

$$OZ(0,0) \ a_{A}^{O} \ a_{B}^{O} + OZ(0,1) \ a_{A}^{O} \ a_{B}^{1} + Z(1,0) \ a_{A} \ a_{B}^{O} +$$

$$C_{A} = \frac{\sum_{j} i \ Z(i,j) \ a_{A}^{i} \ a_{B}^{j}}{Z(0,0) \ a_{A}^{O} a_{B}^{O} + Z(1,0) \ a_{A}^{1} a_{B}^{O} + Z(0,1) \ a_{A}^{O} a_{B}^{1} +}$$

$$\sum_{j} \sum_{i} Z(i,j) \ a_{A}^{i} \ a_{B}^{j}$$

or

$$C_{A} = \frac{Z(1,0) \ a_{A} + \sum \ i \ Z(i,j) \ a_{A}^{i} \ a_{B}^{j}}{Z(0,0) + Z(1,0) \ a_{A} + Z(0,1) \ a_{B}^{+} \sum \ \sum Z(i,j) \ a_{A}^{i} \ a_{B}^{j}}$$

$$j \ i \ (5.23)$$

where summations are now carried out overall values of i and j satisfying

$$i + j \ge 2 \tag{5.24}$$

$$i\beta_A + j\beta_B \leq v$$
 (5.11)

Substitution of eqs.(5.20) to (5.22) in eq.(5.23) gives

$$C_{A} = \frac{\alpha_{A}p_{A} + \sum \sum i Z(i,j) a_{A}^{i} a_{B}^{j}}{1 + \alpha_{A}p_{A} + \alpha_{B}p_{B} + \sum \sum Z(i,j) a_{A}^{i} a_{B}^{j}}$$
(5.25)

Equations (5.17) and (5.25) give
$$\frac{iZ(1,0)^{\dot{1}} a_{A}^{\dot{1}} Z(0,1)^{\dot{j}} a_{B}^{\dot{j}}}{iZ(1,0)^{\dot{1}} a_{\dot{1}}^{\dot{1}} Z(0,1)^{\dot{1}} a_{\dot{1}}^{\dot{1}}} (1-i\beta_{\dot{1}}/v-j\beta_{\dot{1}}/v)^{\dot{1}+\dot{1}} \times \\ \exp \left(\frac{i\beta_{\dot{1}} \epsilon_{\dot{1}} + j\beta_{\dot{1}} \epsilon_{\dot{1}}}{v k T}\right)$$

$$\frac{\sum_{A} = \frac{\sum_{A} \sum_{A} \sum_{$$

Substitution of eqs.(5.21) and (5.22) in above equation gives

$$c_{A} = \frac{\alpha_{A}p_{A} + \sum_{j} \sum_{i} \frac{(\alpha_{A}p_{A})^{i} (\alpha_{B}p_{B})^{j}}{(i-1)! j!} (1-i\beta_{A}/v - j\beta_{B}/v)^{i+j} \times (1-i\beta_{A}/v - j\beta_{B}/v)^{i+j} \times (1-i\beta_{A}/v - j\beta_{B}/v)^{i+j} \times (1-i\beta_{A}/v - j\beta_{B}/v)^{i+j} \times (1-i\beta_{A}p_{A} + \alpha_{B}p_{B} + \sum_{j} \sum_{i} \frac{(\alpha_{A}p_{A})^{i} (\alpha_{B}p_{B})^{j}}{i! j!} (1-i\beta_{A}/v - j\beta_{B}/v)^{i+j} \times (1-i\beta_{A}p_{A} + \alpha_{B}p_{B} + \sum_{j} \sum_{i} \frac{(\alpha_{A}p_{A})^{i} (\alpha_{B}p_{B})^{j}}{i! j!} (1-i\beta_{A}/v - j\beta_{B}/v)^{i+j} \times (1-i\beta_{A}p_{A} + \alpha_{B}p_{B} + \sum_{j} \sum_{i} \frac{(\alpha_{A}p_{A})^{i} (\alpha_{B}p_{B})^{j}}{i! j!} (1-i\beta_{A}/v - j\beta_{B}/v)^{i+j} \times (1-i\beta_{A}p_{A} + \alpha_{B}p_{B} + \sum_{j} \sum_{i} \frac{(\alpha_{A}p_{A})^{i} (\alpha_{B}p_{B})^{j}}{i! j!} (1-i\beta_{A}/v - j\beta_{B}/v)^{i+j} \times (1-i\beta_{A}p_{A} + \alpha_{B}p_{B} + \sum_{j} \sum_{i} \frac{(\alpha_{A}p_{A})^{i} (\alpha_{B}p_{B})^{j}}{i! j!} (1-i\beta_{A}/v - j\beta_{B}/v)^{i+j} \times (1-i\beta_{A}p_{A} + \alpha_{B}p_{B} + \sum_{j} \sum_{i} \frac{(\alpha_{A}p_{A})^{i} (\alpha_{B}p_{B})^{j}}{i! j!} (1-i\beta_{A}/v - j\beta_{B}/v)^{i+j} \times (1-i\beta_{A}p_{A} + \alpha_{B}p_{B} + \sum_{j} \sum_{i} \frac{(\alpha_{A}p_{A})^{i} (\alpha_{B}p_{B})^{j}}{i! j!} (1-i\beta_{A}/v - j\beta_{B}/v)^{i+j} \times (1-i\beta_{A}p_{A} + \alpha_{B}p_{B} + \sum_{j} \sum_{i} \frac{(\alpha_{A}p_{A})^{i} (\alpha_{B}p_{B})^{j}}{i! j!} (1-i\beta_{A}/v - j\beta_{B}/v)^{i+j} \times (1-i\beta_{A}/v - j\beta_{B}/v)^{$$

Similarly equation for $C_{\overline{B}}$ can also be derived, which is

$$\alpha_{B}p_{B} + \sum_{j} \sum_{i} \frac{(\alpha_{A}p_{A})^{i}(\alpha_{B}p_{B})^{j}}{i! (j-1)!} (1-i\beta_{A}/v-j\beta_{B}/v)^{i+j} \times$$

$$= \exp(\frac{i\beta_{A}}{v} \frac{\varepsilon_{A}}{k} + \frac{j\beta_{B}}{k} \frac{\varepsilon_{B}}{k})$$

$$C_{B} = \frac{1+\alpha_{A}p_{A} + \alpha_{B}p_{B} + \sum_{j} \sum_{i} \frac{(\alpha_{A}p_{A})^{i}(\alpha_{B}p_{B})^{j}}{i! j!} (1-i\beta_{A}/v-j\beta_{B}/v)^{i+j} \times$$

$$= \exp(\frac{i\beta_{A}}{v} \frac{\varepsilon_{A}}{k} + \frac{j\beta_{B}}{k} \frac{\varepsilon_{B}}{k})$$

$$= \exp(\frac{i\beta_{A}}{v} \frac{\varepsilon_{A}}{k} + \frac{j\beta_{B}}{k} \frac{\varepsilon_{B}}{k})$$

$$= \exp(\frac{i\beta_{A}}{v} \frac{\varepsilon_{A}}{k} + \frac{j\beta_{B}}{k} \frac{\varepsilon_{B}}{k})$$

The expressions along with constraints (5.11) and (5.24) give the general model for binary mixtures. As in pure component model, when exponential terms are neglected, we get

$$\alpha_{A}p_{A} + \sum_{j} \frac{(\alpha_{A}p_{A})^{i}(\alpha_{B}p_{B})^{j}}{(i-1)! j!} (1-i\beta_{A}/v-j\beta_{B}/v)^{i+j}$$

$$C_{A} = \frac{1}{1+\alpha_{A}p_{A}+\alpha_{B}p_{B}} + \sum_{j} \frac{(\alpha_{A}p_{A})^{i}(\alpha_{B}p_{B})^{j}}{i! j!} (1-i\beta_{A}/v - j\beta_{B}/v)^{i+j}$$

$$(5.29)$$

and

$$C_{B} = \frac{\alpha_{A}p_{B} + \sum_{j} \sum_{i} \frac{(\alpha_{A}p_{A})^{i}(\alpha_{B}p_{B})^{j}}{i! (j-1)!} (1-i\beta_{A}/v - j\beta_{B}/v)^{i+j}}{1+\alpha_{A}p_{A} + \alpha_{B}p_{B} + \sum_{j} \sum_{i} \frac{(\alpha_{A}p_{A})^{i}(\alpha_{B}p_{B})^{j}}{i! j!} (1-i\beta_{A}/v-j\beta_{B}/v)^{i+j}}$$
(5.30)

Equations (5.29) and (5.30) along with constraints (5.11) and (5.24) give simplified form of the model for binary mixtures. Values of α_A , α_B , β_A and β_B needed in above expressions are obtained from isotherms of pure components. Raoult's law for components A and B gives

$$p_A = P_A x_A$$
 and $p_B = P_B x_B$

where P_A and P_B are vapor pressures of components A and B respectively and \mathbf{x}_A and \mathbf{x}_B are molefractions of A and B in liquid phase. Substitution of these in equations (5.29) and (5.30) gives

$$c_{A} = \frac{\alpha_{A}P_{A}x_{A} + \sum_{j} \sum_{i} \frac{(\alpha_{A}P_{A}x_{A})^{i}(\alpha_{B}P_{B}x_{B})^{j}}{(i-1)! j!} (1-i\beta_{A}/v -j\beta_{B}/v)^{i+j}}{1+\alpha_{A}P_{A}x_{A} + \alpha_{B}P_{B}x_{B} + \sum_{j} \sum_{i} \frac{(\alpha_{A}P_{A}x_{A})^{i}(\alpha_{B}P_{B}x_{B})^{j}}{i! j!} (1-i\beta_{A}/v-j\beta_{B}/v)^{i+j}}$$
(5.31)

and

$$C_{B} = \frac{\alpha_{B}P_{B}x_{B} + \sum_{j} \sum_{i} \frac{(\alpha_{A}P_{A}x_{A})^{i}(\alpha_{B}P_{B}x_{B})^{j}}{i!(j-1)!} (1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v})^{i+j}}{1 + \alpha_{A}P_{A}x_{A} + \alpha_{B}P_{B}x_{B} + \sum_{j} \sum_{i} \frac{(\alpha_{A}P_{A}x_{A})^{i}(\alpha_{B}P_{B}x_{B})^{j}}{i!j!} x}$$

$$(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v})^{i+j} \qquad (5.32)$$

Equations (5.31) and (5.32) along with the constraints (5.11) and (5.24) were used to calculate average number of molecules of A and B adsorbed per cavity in a binary system of A and B. Values of α_A , α_B , β_A and β_B needed in above expressions are obtained from isotherms of pure components.

5.4 EXTENSION OF THE MODEL TO TERNARY SYSTEMS:

The above model can be extended to a multicomponent system with any number of components. We extend it here for three components. In this case, there are three components A, B and C with Henry's constants α_A , α_B and α_C and molecular volumes β_A , β_B and β_C respectively. The grand partition function, G_{ABC} , for the system can be written as

$$G_{ABC} = \begin{bmatrix} \sum \sum \sum q(i,j,h) & \lambda_A^i & \lambda_B^j & \lambda_C^h \end{bmatrix}^M = Q_{ABC}^M$$

with constraint

$$i\beta_A + j\beta_B + h\beta_C \leq v$$

where q(i,j,h) is the partition function of a cavity containing i molecules of A, j molecules of B and h molecules of C. Q_{ABC} is the canonical ensemble partition function for the system containing components A, B and C. $\lambda_A=e^{\mu_A/kT}$, $\lambda_B=e^{\mu_B/kT}$ and $\lambda_C=e^{\mu_C/kT}$ (μ_A , μ_B and μ_C are chemical potentials for components A, B and C respectively). In terms of configuration integrals above equation can be written as

$$G_{ABC} = \begin{bmatrix} \sum_{h} \sum_{j} \sum_{i} Z(i,j,h) a_{A}^{i} a_{B}^{j} a_{C}^{h} \end{bmatrix} = Q_{ABC}^{M}$$
 (5.33)

where Z(i,j,h) is the configuration integral for a cavity containing i molecules of A, j molecules of B and h molecules of C. a_A , a_B and a_C are activities of A, B and C respectively. The average number of molecules of A, B and C adsorbed can be written as

$$C_A = \frac{n_A}{M} = \lambda_A \left(\frac{\partial \ln Q_{ABC}}{\partial \lambda_A} \right) \lambda_B, \lambda_C, T$$
 (5.34)

$$C_B = \frac{n_B}{M} = \lambda_B \left(\frac{\partial \ln Q_{ABC}}{\partial \lambda_B} \right) \lambda_A, \lambda_C, T$$
 (5.35)

$$C_C = \frac{n_C}{M} = \lambda_C(\frac{3 \ln Q_{ABC}}{3 \lambda_C}) \lambda_A, \lambda_B, T$$
 (5.36)

Equation (5.33) with equations (5.34 to 5.35) gives

$$C_{A} = \frac{\sum_{h} \sum_{j} \sum_{i} Z(i,j,h) a_{A}^{i} a_{B}^{j} a_{C}^{h}}{\sum_{h} \sum_{j} \sum_{i} Z(i,j,h) a_{A}^{i} a_{B}^{j} a_{C}^{h}}$$
(5.37)

$$C_{B} = \frac{\sum_{h} \sum_{j} \sum_{j} Z(i,j,h) a_{A}^{i} a_{B}^{j} a_{C}^{h}}{\sum_{h} \sum_{j} \sum_{j} Z(i,j,h) a_{A}^{i} a_{B}^{j} a_{C}^{h}}$$
(5.38)

$$C_{C} = \frac{\sum_{A} \sum_{A} \sum_{A}$$

The configuration integrals for ternary systems, with the usual assumptions as discussed earlier, are approximated by

$$Z(i,j,h) = \begin{bmatrix} \frac{Z(1,0,0)^{i} Z(0,1,0)^{j} Z(0,0,1)^{h}}{i! j! h!} (1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h} \\ \exp \left(\frac{i\beta_{A}\epsilon_{A} + j\beta_{B}\epsilon_{B} + h\beta_{C}\epsilon_{C}}{v k T}\right) \end{bmatrix}$$
(5.40)

where.

$$Z(0,0,0) = Z(0) = 1$$
 (5.41)

$$Z(1,0,0) a_A = Z_A a_A = \alpha_A p_A$$
 (5.42)

$$Z(0,1,0) a_B = Z_B a_B = \alpha_B p_B$$
 (5.43)

$$Z(0,0,1) a_C = Z_C a_C = \alpha_C p_C$$
 (5.44)

where p_A , p_B and p_C are partial pressures of components A,B and C respectively. Equation (5.37) on expansion gives

$$C_{A} = \frac{a_{A}^{0}a_{B}^{0}a_{C}^{0} + 0Z(0,1,0) \ a_{A}^{0}a_{B}^{1}a_{C}^{0} + 0Z(0,0,1) \times}{a_{A}^{0}a_{B}^{0}a_{C}^{1} + 1Z(1,0,0) \ a_{A}^{1}a_{B}^{0}a_{C}^{0} + \sum_{h=1}^{2} \sum_{i=1}^{4} iZ(i,j,h) \times}$$

$$C_{A} = \frac{a_{A}^{i} \ a_{B}^{j} \ a_{C}^{h}}{Z(0,0,0) \ a_{A}^{0} \ a_{B}^{0} \ a_{C}^{0} + Z(1,0,0) \ a_{A}^{1} \ a_{B}^{0}a_{C}^{0} + Z(0,1,0) \times}$$

$$a_{A}^{0}a_{B}^{1}a_{C}^{0} + Z(0,0,1) \ a_{A}^{0}a_{B}^{0}a_{C}^{1} + \sum_{h=j=1}^{2} \sum_{i=1}^{2} Z(i,j,h) \times}$$

$$a_{A}^{i} \ a_{B}^{j} \ a_{C}^{h}$$

with a added constraint i+j+h ≥ 2

Equations (5.41) to (5.44) with above equation gives

$$C_{A} = \frac{\alpha_{A} p_{A} + \sum\limits_{h} \sum\limits_{j} \sum\limits_{i} iZ(i,j,h), a_{A}^{i} a_{B}^{j} a_{C}^{h}}{1 + \alpha_{A} p_{A} + \alpha_{B} p_{B} + \alpha_{C} p_{C} + \sum\limits_{h} \sum\limits_{j} \sum\limits_{i} Z(i,j,h) a_{A}^{i} a_{B}^{j} a_{C}^{h}}$$

Substitution of equation (5.40) in above equation gives

$$\alpha_{A}p_{A} + \sum_{h j i} \sum_{i} \frac{z(1,0,0)^{i} a_{A}^{i} z(0,1,0)^{j} a_{B}^{j} z(0,0,1)^{h} a_{C}^{h}}{(i-1)! j! h!} \times \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h} \exp(\frac{i\beta_{A} \epsilon_{A} + j\beta_{B} \epsilon_{B} + h\beta_{C} \epsilon_{C}}{v k T})$$

$$C_{A} = \frac{1+\alpha_{A}p_{A}+\alpha_{B}p_{B}+\alpha_{C}p_{C}}{1+\alpha_{A}p_{A}+\alpha_{B}p_{B}+\alpha_{C}p_{C}} + \sum_{h j i} \sum_{i} \frac{z(1,0,0)^{i}a_{A}^{i} z(0,1,0)^{j}a_{B}^{j} z(0,0,1)^{h}}{i! j! h!}$$

$$(1-\frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h} \exp(\frac{i\beta_{A}\epsilon_{A} + j\beta_{B}\epsilon_{B} + h\beta_{C}\epsilon_{C}}{v k T})$$

Equations (5.42) to (5.44) with above equation gives

$$\alpha_{A}p_{A} + \sum_{h} \sum_{j} \sum_{i} \frac{(\alpha_{A}p_{A})^{i}(\alpha_{B}p_{B})^{j}(\alpha_{C}p_{C})^{h}}{(i-1)!j!h!} \times$$

$$(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h} \exp(\frac{i\beta_{A}\epsilon_{A} + j\beta_{B}\epsilon_{B} + h\beta_{C}\epsilon_{C}}{v k T})$$

$$C_{A} = \frac{1 + \alpha_{A}p_{A} + \alpha_{B}p_{B} + \alpha_{C}p_{C} + \sum_{h} \sum_{j} \sum_{i} \frac{(\alpha_{A}p_{A})^{i}(\alpha_{B}p_{B})^{j}(\alpha_{C}p_{C})^{h}}{i!j!h!} \times$$

$$(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h} \exp(\frac{i\beta_{A}\epsilon_{A} + j\beta_{B}\epsilon_{B} + h\beta_{C}\epsilon_{C}}{v k T})$$

$$(5.45)$$

With constraints $i+j+h \ge 2$ and $i\beta_A + j\beta_B + h\beta_C \le v$.

Equation (5.45) gives the molecules of component A adsorbed per cavity of zeolite. Similar expressions for components B and C can be derived and are

$$\alpha_{B}p_{B} + \sum_{h} \sum_{j} \sum_{i} \frac{(\alpha_{A}p_{A})^{i}(\alpha_{B}p_{B})^{j}(\alpha_{C}p_{C})^{h}}{i! (j-1)! h!} \times$$

$$C_{B} = \frac{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h} \exp(\frac{i\beta_{A}\epsilon_{A} + j\beta_{B}\epsilon_{B} + h\beta_{C}\epsilon_{C}}{v k T})}{1 + \alpha_{A}p_{A} + \alpha_{B}p_{B} + \alpha_{C}p_{C} + \sum_{h} \sum_{j} \frac{(\alpha_{A}p_{A})^{i}(\alpha_{B}p_{B})^{j}(\alpha_{C}p_{C})^{h}}{i! j! h!} \times (1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h} \exp(\frac{i\beta_{A}\epsilon_{A} + j\beta_{B}\epsilon_{B} + h\beta_{C}\epsilon_{C}}{v k T})}{(5.46)}$$

$$C_{C} = \frac{1 + \alpha_{A} p_{A} + \alpha_{B} p_{B} + \alpha_{C} p_{C} + \sum_{h \ j \ i} \sum_{i} \frac{(\alpha_{A} p_{A})^{i} (\alpha_{B} p_{B})^{j} (\alpha_{C} p_{C})^{h}}{i! \ j! \ h!} \times \frac{1 + \alpha_{A} p_{A} + \alpha_{B} p_{B} + \alpha_{C} p_{C}}{v} + \sum_{h \ j \ i} \sum_{i} \frac{(\alpha_{A} p_{A})^{i} (\alpha_{B} p_{B})^{j} (\alpha_{C} p_{C})^{h}}{i! \ j! \ h!} \times \frac{1 + \alpha_{A} p_{A} + \alpha_{B} p_{B} + \alpha_{C} p_{C}}{v \ k \ T}$$

$$(5.47)$$

with constraints

$$i\beta_{A} + j\beta_{B} + h\beta_{C} \leq v$$

$$i+j+h \geq 2$$

$$(5.48)$$

Equations (5.45) to (5.47) along with the constraints (5.48) and (5.49) predict the value of molecules adsorbed per cavity for components A, B and C respectively in a ternary system. When the exponential term representing sorbate-sorbate attraction is neglected, the above equations reduce to

$$\begin{split} c_{A} &= \frac{\alpha_{A} p_{A} + \sum\limits_{h} \sum\limits_{j} \sum\limits_{i} \frac{(\alpha_{A} p_{A})^{i} (\alpha_{B} p_{B})^{j} (\alpha_{C} p_{C})^{h}}{(i-1)! j! h!} (1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h}} \\ c_{A} &= \frac{1 + \alpha_{A} p_{A} + \alpha_{B} p_{B} + \alpha_{C} p_{C} + \sum\limits_{h} \sum\limits_{j} \sum\limits_{i} \frac{(\alpha_{A} p_{A})^{i} (\alpha_{B} p_{B})^{j} (\alpha_{C} p_{C})^{h}}{i! j! h!} \times \\ & \cdot (1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h} & (5.50) \\ c_{B} &= \frac{\alpha_{B} p_{B} + \sum\limits_{h} \sum\limits_{j} \sum\limits_{i} \frac{(\alpha_{A} p_{A})^{i} (\alpha_{B} p_{B})^{j} (\alpha_{C} p_{C})^{h}}{i! (j-1)! h!} (1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h} \\ c_{B} &= \frac{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h}}{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h}} & (5.51) \\ c_{C} &= \frac{\alpha_{C} p_{C} + \sum\limits_{h} \sum\limits_{j} \sum\limits_{i} \frac{(\alpha_{A} p_{A})^{i} (\alpha_{B} p_{B})^{j} (\alpha_{C} p_{C})^{h}}{i! j! (h-1)!} (1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h}} \\ c_{C} &= \frac{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h}}{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h}} & (5.52) \\ c_{C} &= \frac{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h}}{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h}} & (5.52) \\ c_{C} &= \frac{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h}}{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h}} & (5.52) \\ c_{C} &= \frac{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h}}{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h}} & (5.52) \\ c_{C} &= \frac{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h}}{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h}} & (5.52) \\ c_{C} &= \frac{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h}}{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h}} & (5.52) \\ c_{C} &= \frac{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h}}{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h}} & (5.52) \\ c_{C} &= \frac{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{i+j+h}}{(1 - \frac{i\beta_{C}}{v} - \frac{j\beta_{C}}{v} - \frac{j\beta_{C}}{v} - \frac{j\beta_{C}}{v} - \frac{j\beta_{C}}{v} - \frac{j\beta_{C}}{v} - \frac{j\beta_{C}}{v} - \frac{j\beta_{C}$$

Using Raoult's law to calculate the partial pressures, one gets

$$C_{A} = \frac{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{1} + j + h}{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{1} + j + h} \times (5.55)$$

$$C_{B} = \frac{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{1} + j + h}{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{1} + j + h} \times (5.55)$$

$$C_{B} = \frac{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{1} + j + h}{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{1} + j + h} \times (5.55)$$

$$C_{C} = \frac{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{1} + j + h}{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{1} + j + h} \times (5.54)$$

$$C_{C} = \frac{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{1} + j + h}{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{1} + j + h} \times (5.54)$$

$$C_{C} = \frac{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{1} + j + h}{(1 - \frac{i\beta_{A}}{v} - \frac{j\beta_{B}}{v} - \frac{h\beta_{C}}{v})^{1} + j + h} \times (5.55)$$

where x_A , x_B and x_C are mole fractions of A, B and C in liquid phase and P_A , P_B and P_C are vapor pressures of components A, B and C respectively. Equations (5.53), (5.54)

and 5.55 along with the constraints(5.48) and (5.49) were used to calculate average number of molecules adsorbed per cavity of components A, B and C respectively. α_A , β_A , α_B , β_B , α_C and β_C used in these equations were obtained from isotherms of pure components.

5.5 RESULTS AND DISCUSSION:

5.5.1 Adsorption of Pure Hydrocarbons:

The equation derived from the model for isotherms of pure component (Eq. (5.9)) has two parameters α and β. α can be calculated from the slope of the adsorption isotherm at very low concentrations. The nature of the isotherms (Figures 3. 3to 3.6) did not permit direct measurement of the slope, because of experimental limitations of finding equilibrium loadings at very low concentrations. It was also observed that, for the n-paraffins studied, equation (5.9) is very sensitive to eta_{ullet} the molecular volume of the adsorbate. approximation of its value by van der Waals co-volume, as suggested by Dubinin and used by Ruthven and coworkers [14-18]. was found to be unsatisfactory for the systems investigated. Thus it is observed that a more accurate determination of the values of the parameters α and β is required to predict the adsorption isotherms for the hydrocarbons studied. However, in the absence of better prediction procedures for α and β , it was decided to obtain these using the experimental

data. Thus, it has been implicitly assumed that our two parameter model is valid for pure hydrocarbon adsorption. Experimental isotherms (Data are taken from A3.3 to A3.6) were matched with the model by a least square minimization procedure, using Blind search Technique to evaluate α and β . An objective function E is defined as

$$E = \begin{bmatrix} \sum_{i=1}^{n} (c_{cal} - c_{Expt})^2 \end{bmatrix}$$

where, n = number of data points

C_{Cal} = calculated value of molecules adsorbed per cavity from Eq.(5.9)

 $C_{\rm Expt}$ = experimental value of molecules adsorbed per cavity E is then minimized over a wide range of the values of parameters α and β to give minimum value, $E_{\rm min}$, of the objective function.

$$E_{\text{Min}} = \frac{\text{Min}}{\alpha, \beta} \left[\sum_{i=1}^{n} (c_{\text{Cal}} - c_{\text{Expt}})^2 \right]$$

The values of α and β corresponding to E_{min} were then taken to represent the best values of these parameters for a given paraffin at a fixed temperature. Figure 5.1 shows typical predicted adsorption isotherms using calculated values of α and β along with experimentally observed data points at $18^{\circ}C$. Similar plots were obtained at other temperatures also but have not been included here. This

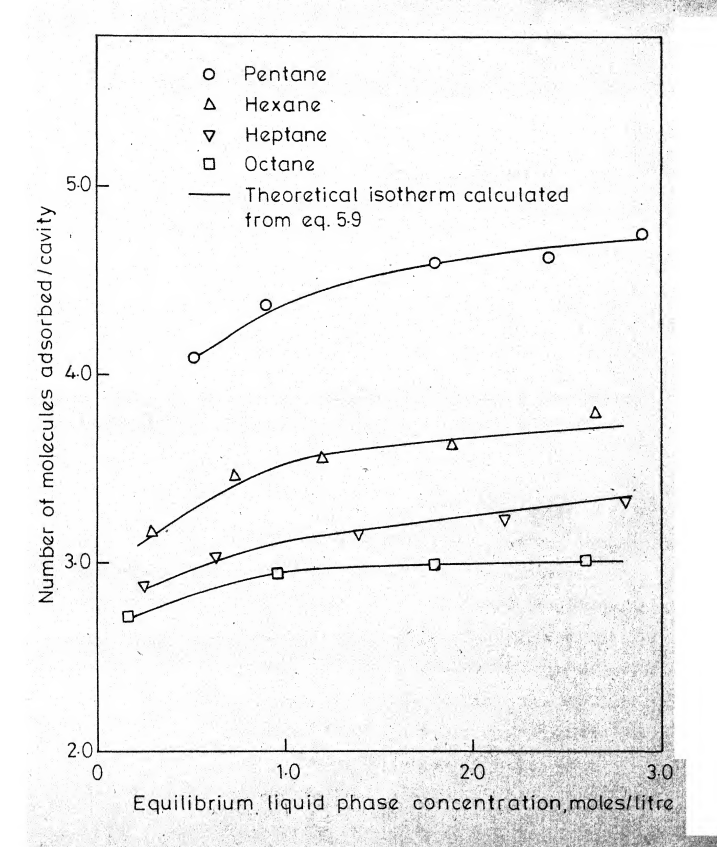


Fig. 5.1 - Comparison between predicted and experimental isotherms at 18°C.

should not be taken as a validation of the model because the prediction is not totally independent of experimental data, rather, it makes use of the latter to calculate the former. The comparison has been made to merely show that the two parameter model is able to represent the experimental data fairly well. As will be seen later, the validation of the pure component model comes from the fact that, for most of the systems studied, the predictions made using this model for binary and ternary systems match within 10 per cent of the experimentally observed values.

The values of α and β along with the minimum error, for all the paraffins studied, at different temperatures are given in Table 5.1. α was found to decrease with an increase in temperature which means that adsorption decreases with increasing temperature. It was also observed that α increased with the molecular weight of paraffin. The molecular volume of the adsorbate, as expected, increased with temperature and was higher, at a particular temperature, for higher molecular weight paraffin. The internal consistency in calculated values of α and β further justifies the way they were obtained. Figure 5.2 shows Arrhenius plots for all the hydrocarbons studied.

TABLE 5.1: VALUES OF & AND 8 AT DIFFERENT TEMPERATURES

Paraffin	Temperature,	$\alpha_{\it j}$ (molecules)/(cavity)(torr)	β, A3	Emin
n-pentane	6 1.8 30	4.50 3.96 1.86	94.50 116.55 129.50	0.0521
n-hexane	6 18 30 42	64.05 50.92 46.03 22.46	136.50 165.71 179.88 183.79	0.0121 0.0241 0.0465
n-heptane	18 30 42	222.43 222.43 101.44 36.14	166.19 179.31 183.96 184.99	0.0227 0.0199 0.0488 0.0877
n-Octane	18 30 42	3079.89 824.36 485.90 247.75	181.92 187.41 215.00 225.10	0.0019

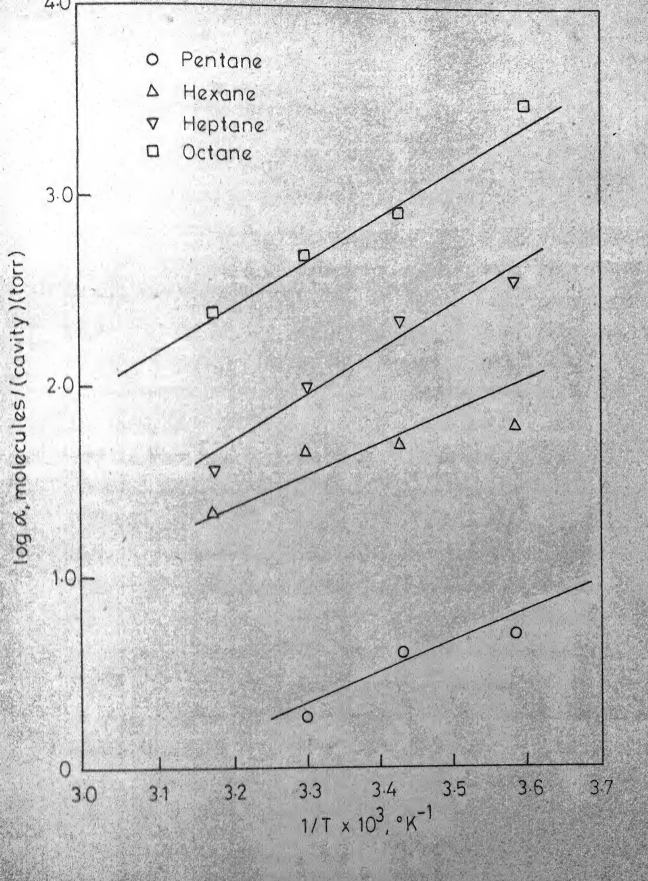


Fig 5 2 - Arrhenius plot for &.

5.5.2 Adsorption of Binary Systems:

Equations (5.31) and (5.32) along with constraints (5.11) and (5.24) were used to predict values of C_A and C_B , for all the binary systems studied, using appropriate values of α and β obtained for pure components. Tables 5.2 to 5.5 (operating data are given in A4.1 to A4.4) give the results of all the systems at 6°, 18°, 30° and 42°C respectively. In general errors are less than 10 per cent, for more than 90 per cent of a total of 136 predictions made. At 6°C, the maximum error of 10.8 per cent is observed in the case of pentane-hexane system, all others out of a total of 40 predictio showing less than 10 per cent error. At 18°C, the maximum error of 12.4 per cent is also observed for the pentane-hexane system. There are 3 other predicted values which are more than 10 per cent higher than the experimental values at this temperature. The difference in predicted and experimental value of molecules adsorbed per cavity increases upto about 14 per cent, at 30°C, for systems having pentane as one component. This large error in case of pentane at 30°C may be attributed to larger thermal motions of pentane molecules adsorbed at this temperature which is close to its boiling point. Due to these motions some of the adsorbed pentane may come out of the zeolite cavity resulting in smaller loading than predicted. The proposed model does not account for any such desorption and hence predicts higher adsorption than

TABLE 5.2: COMPARISON OF EXPERIEBULAL AND PREDICTED VALUES FOR BINARY SYSTEMS AT 6

	Equilibrium	composition in	Molecules	les adsorbed	per	cavi ty	Per cent,	Erro
Run	liquid phase	liquid phase, mole fraction	Experimental	mental	Pred	Predicted		t
	1	2	1	2	٦	2		
BPX-1	0.321	0.052	4.75	1,36	4.88	1.32	2.7	2.5
BPX-2	0.213	0.115	3.20	2.01	3.44	2.00	7.3	0*3
BPX-3	0.134	0.201	1.96	2.98	2,17	* 2.95	10,8	1.0
BPX-4	0.052	0.290	1.31	3.90	~	. •	1.7	0.5
BXH-1	0.063	0.284	1.36	2,91	1.41	2.95	3.6	1.4
BXH-2	0.138	0.214	2.02	2,13	2.06	2.17	1,9	2.0
BXH-3	0.206	0.144	2.75	1.59	2.81	1.64	2.0	3.3
BXH-4	0.285	0.062	3.69		3.78	1.21	2.3	3.4
BH0-1	0.055	0.278	1.17	2,89	1.25	2.92	L L L 9	1.1
BH0-2	0.127	0.198	1.61	2,24	1.69	2,30	5.4	2.7
BH0-3	0.208	0.122	2.31	1.61	2.35	1.70	1.6	5.7
BH0-4	0.280	0.053	3.04	1,20	3,11	1.27	2.2	6.1
1			1	1	1		1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	1 1 1 1

BPX- Pentane-heXane
BXH= heXane-Heptane
BHO=Heptane-Octane
BPH= Pentane-Heptane
BPO= Pentane-Octane

Table 5.2 (contd)

Bun	Equilibrium composition	omposition in	Molecule	s adsorbe	Molecules adsorbed per cavity	i ty	Per cent, Error	rror
	liquid phase	liquid phase mole Traction	haperimentar 1	2	I	2		2
BPH-1	0.302	0.042	5.00	1,13	5.14	1.16	2.7	2.4
BPH-2	0.237	0.094	3.92	1.47	4.15	1.45	5.9	1.3
BPH-3	0.157	0.164	2.62	2,11	2,85	2.04	0.6	3.4
BPH-4	0.065	0.263	1.44	3.10	1.57	3.05	0.6	1.4
			1 1 1		1 1 1		1 1 1 1	1 0
BP0-1	0.054	0.275	1.20	2.94	1,28	2.93	7.	0.0
BP0-2	0.141	0.203	2.05	3.22	2.17	3.29	5.7	2.2
BP0-3	0.219	711.0	3.27	1.59	3.44	1.65	5.2	3.9
BP0-4	0.314	0.052	4.77	1.11	4.85	1.21	∞ . ⊢	9.1
Note:	l represents the lighter remaining component.	1	hydrocarbon in a given binary mixture	in a give	n binary	mixture	and 2 represents	nts the

TABLE 5.3: COMPARISON OF EXPERINGNTAL AND PREDICTED VALUES FOR BINARY SYSTEMS AT 1800

Run	Equilibrium composition liquid phase, mole frac	composition in mole fraction 2	Molecules Experim	lecules adsorbed Experimental 1 2		er cavity Predicted 1 2	Per cent,	Frror 2
BPX-5	0.327	0.052	4.05	1.07	4.17	1,20	3.0	12.4
BPX-6	0.231	0.119	3.04	1.55	3.14	1.64	3.2	5.4
BPX-7	0.152	0.203	1.99	2,23	2,11	2,32	6.2	4.2
BPX-8	0.064	0.288	1.22	3.10	1.36	3.19	11.6	2.9
I	1 1 90 0		1 90 1	1 0	1 6	1 0	1 5	_
, - mwa	† 00 · 0	0.23.0		00.4	1.73	7	FO. 4	T.C
BXH-6	0.146	0.201	1.84	1.90	2.01	1,98	9.1	4.1
BXH-7	0.219	0.130	2,51	1.46	2.63	1.52	4.7	4.4
BXH-8	0.302	0.061	3.14	1.13	3.29	1.20	4.9	6.4
BHO-5	0.052	0.248	1.16	2.49	1.23	2.52	6.4	1 1 1
9-0HB	0.114	991.0	1.57	1.85	1,65	1,93	5.3	4.0
BH0-7	0.178	0.108	2.03	1.42	2,17	1.50	8.9	5.6
BH0-8	0.264	0.045	2,82	1.08	2,89	1.16	2.3	7.2

Table 5.3 (contd)

Run	Equilibrium composition liquid phase, mole frac	Equilibrium composition in liquid phase, mole fraction 1	Molecules adsorbed per cavity Experimental Predicted 1	adsorbed tal	per cavit Predicted	ity ed 2	Per cent,	Brror 2
BPH-5	0.320	0.045	4.13	1.07	4.29	1.14	3.9	6.2
BPH-6	0.243	0.098	3.39	1.35	3.45	1.42	1.9	5.5
BPH-7	0.177	0.168	2,36	1.82	2.51	1,88	6.4	3.7
BPH-8	0.070	0.264	1,29	2,63	•	2.74	10.2	4.1
BP0-5	790.0	0.249	1.36	2.54	1.48	2.56	0.6	
BP0-6	0.156	0.170	2,25	1.81	2,38	1.89	5.7	4.4
BP0-7	0.237	0.114	3.10	1,35	3.32	1.45	7.0	7.2
BP0-8	0.327	0.045	4.26	1.05	4.33	1.12	1.7	8*9
			L					

ote: Notations are same as used in Table 5.2

TABLE 5.4: COMPANISON OF BAPERINENTAL AND PREDICTED VALUES FOR BINARY SYSTEMS AT 300

	The state of the s								
Run	Equilibrium composition liquid phase, mole frac	position in ole fraction 2	Molecules adsc Experimental	s adsorbed lental 2	per Predi 1	cavity	Per cent,	Brror 2	1 .
BPX-9	0.074	0.288	1.04	2.80	1,19	3.06	14.4	9.3	1
BPX-10	0.170	0.205	1.42	2.39	1.56	2.54	6.6	2.9	
BPX-11	0.247	0.123	1.88	1.92	2.09	2.04	10.9	0*9	
BPX-12	0.335	0.052	2.96	1.34	3.13	1.44	5.6	7.6	
1 1			1 1 1		1				
BXH-9	0.317	0.050	3.04	1.04	3,14	1,10	3.1	6.0	1
BXH-10	0.251	0.135	2.54	1,24	2,62	1,32	K.	5,9	
BXH-11	0.155	0.189	1.98	1.55	2.07	1.62	4.2	4.5	
BXH-12	0.064	0.274	1.33	2.20	1.41	2,30	6.2	4.	•
BH0-9	0.049	0.280		1 72 0	1 6	1 1		1 1 1	
RHO_10	0		1	07.7	1.60	Z + 4 T	T•)	2.5	
0T-0HG	0.104	0.197	1.43	1.81	1.52	1,88	6.4	3.8	
BH0-11	0.180	0.119	1.93	3.53	2.06	1.42	6.5	۲. ۲	
BH0-12	0.270	0.050	2.55	1.06	2.68	1,14		7.7	
1 1 1			1 1 1	1	1	1	1 1 1 1	1 1 1	_

Table 5.4 (contd)

Run	Equilibrium composition in liquid phase, mole fraction 2	mposition in mole fraction 2	Molecules adsor Experimental		adsorbed per cavity ental Predicted 2 1 2	er cavity Predicted	Fer cent,	Brror 2
BPH-9	0.339	0.047	3,38	1,16	3.58	1,24	6.2	7.1
BPH-10	0.268	0.102	2.67	1,49	2.75	1.58	2.8	5.9
BPH-11	0.197	0.171	1,89	1,88	2,06	2,00	8,8	6.1
BPH-12	0.075	0.265	1,20	2.56	1.35	2.70	12,5	5.2
1 1		n eres usus erick king mess wern more			1	-		
BP0-9	0.071	0.279	1,26	2.39	1.44	2.47	13.8	3.2
BP0-10	0.173	0.200	1.96	1.67	2,17	1.78	10.5	6.2
BP0-11	0.254	0.120	2.73	1.29	2.91	1.37	9•9	6.4
BP0-12	0.330	0.051	3.75	1.05	3.73	1.13	0.4	7.4

Notations are same as used in Table 5.2 No te:

TABLE 5.5: COMPARISON OF EXPERINGNTAL AND PREDICTED VALUES FOR BINARY SYSTEMS AT 42°C

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Run	Equilibrium composition liquid phase, mole frac	composition in mole fraction 2	Molecules adsorbed per cavity Experimental Predicted	adsorbertal	d per cavi Predicted	avi ty ted 2	Per cent,	Error 2	
BXH-13	0.063	0.271	1.22	2.01	1,39	2,13	13.5	5.7	1
ВХН-14	0.148	0.193	1,80	1.44	2.01	1.53	11.4	5,8	
BXH-15	0.236	0.120	2.32	1.14	2.52	1.24	8.7	7.8	
BXH-16	0.308	0.050	2,69	0.98	2.92	1,08	8.2	9.2	
BH0-13	0.050	0.275	1,10	2.25	1.20	2.35	8 8	4.3	I
BH0-14	0.120	0.224	1.40	1.78	1.50	1,86	9*9	4.1	
BH0-15	0.190	0.120	1.90	1.34	2,01	1.41	5.5	5.2	
ВНО-16	0.271	0.050	2.49	1.06	2.58	1.14	4.1	7.1	
									ı

Note: Notations are same as used in Table 5.2

that actually taking place. At 42°C, maximum error is found in hexane-heptane system which is 13.5 per cent, for hexane. This can also be justified with the same reasoning as for pentane.

5.5.3 Adsorption of Ternary Systems:

Equations (5.53) to (5.55) along with constraints (5.48) and (5.49) were used to predict values of \mathbf{C}_{A} , \mathbf{C}_{B} and Co, for all the ternary systems studied, using appropriate values of α and β obtained for pure components. Tables 5.6 to 5.8 (Operating conditions are taken from Appendices A4.5 to A4.7) give the results of all the ternary systems studied at 6°. 18° and 30°C respectively. Since pentane was present in all the systems, temperatures higher than 30°C were not studied. It has been observed that for about 85 per cent of a total of 120 predictions made, errors are less than 10 per cent. At 6°C, maximum error is 14 per cent and there are two more predictions for which errors are more than 10 per cent. At 180c, there are 5 predicted values which are more than 10 per cent higher than the experimental values. Errors increase upto 18.5 per cent, at 30°C. There are 11 more predictions at this temperature for which the error is more than 10 per cent. At 30°C, 25 per cent of the predicted values are 10 per cent higher than the experimentally observed values. This does not mean that the ternary

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And in case of the last of the	The second secon			Annual Charles of the Control of the	The same of the sa						
Run	Equiliby liquid	Equilibrium composition	sition in fraction 3	Mole Exper	Molecules Experimenta 1 2	adsorbed 1 3	ed per cavity Predicted	3	Per con	cent, Er	Brror 3
PXH-1	0,235	0.047	0.083	3.29	1.26	1.29	3.68 1.32	1.31	11.9	4.6	1.6
PXH-2	0.082	0.174	0.089	1.45	2,46	1.30	1,66 2,60]	1.33	14.1	5.5	2.3
PXH-3	0.053	0.139	0,165	1,20	2.02	1.72	1.32 2.15]	1,76	6.6	L.9	2.4
PXH-4	0.134	0.087	0.141	1.99	1,61	1.60	2,27 1,66 1	1.61	14.0	3.0	1.0
PX0-1	0.207	0.046	0.045	3.49	1.27	1.15	3.62 1.37 1	-24	3.7	8.5	7.6
PX0-2	0.082	0.149	0.045	1.67	2.50	1.14	1.76 2.59 1	25	5.6	3.4	9.6
PX0-3	0.052	0.132	980.0	1.35	2,26	1.47	1.44 2.31 1	.54	7.0	2.3	5.0
PX0-4	0.124	0.088	0.088	2.11	1.64	1.44	2.22 1.74 1	1.50	5.1	0.9	4.1
PH0-1	0.223	0,056	0.050	3.60	1,16	1,19	3.75 1.23 1	. 25	4.2	5.7	5.2
PH0-2	0.082	980.0	980.0	1.66	1.44	1.62	1.80 1.53 1	1.70	8.5	5.9	4.5
PH0-3	0.017	0,162	0.045	1,32	2.17	1.23	1.45 2.31 1.	.32	7.6	6.5	7.7
PH0-4	0.055	0.130	0.075	1,35	1.83	1.51	1,46 1,91 1	.57	8	4.5	3.7

PHO=Pentane-Heptane-Octane PXO = Pentane-heXane-Octane; PXH = Pentane-heXane-Heptane;

Run	Liquid	rıum phas	position in le fraction	BX	Molecules ad Experimental	adsorbed tal	bed per Predi	er cavity	Per cent.	t. Error	
	7	2	3	7	2	3	7	2 3	-1	1	18
PXH-5	0.114	0.051	0.165	1.66	1,17	1.86	1.80	1.27 1.90	8.4	8.3	2.1
РХН-6	0.119	0.123	0.108	1.62	1,60	1.37	1.79	1.72 1.46	10.3	7.0	6.2
PXH-7	0.216	0.121	0.028	1.74	1.54	0.99	2.85	1.63 1.09	3.9	5.8	10.4
PXH-8	0.129	0.168	0.029	1,85	2.04	0.98	1.96	2.17 1.11	5.8	6.2	12,1
PX0-5	0.045	0.048	0.170	1,22	1.23	1.96	1.33	1.34 2.10	9.1	1 1 8	7.0
PX0-6	0,221	0.073	0.023	3.15	1.31	1.03	3.29	1.41 1.08	4°		• (
PX0-7	0.114	0.144	0.040	1,80	2.04	1.07	1.89	2.09 1.15			7.4
PX0-8	0.056	0.153	0.089	1.21	2.07	1.29	1.34	1.36			5.4
PHO-5	0.101	0.042	0.143	1.70	1.09	1.73	1.86	1.21 1.83	1	1 4	u
PH0-6	0.216	0.044	0.025	3.42	1.11	1.01		21 1.1		ά	
PH0-7	0.063	0.103	0.092	1.33	1.58	1.38	1.51	1.50) . · · ·	• .) α
PH0-8	991.0	660.0	0.036	2.47	1.46	1.06	2.62	1.15	•	•	

Note: Notations are same as used in Table 5.6

	Ramilibrium commosition	m composi	ition in	so [troe [o]]	od conhod	ran ban	1	+17			
E	liquid phe	phase, mole		1451	3	ICAL I	dicted 2	3	Per co	cent, Er	Error 3
РХН-9	0.077	0.221	0.044	1.04 2.54 1	1.04	1.23	2.67	1.11	18.5	5.2	6.4
PXH-10	0.259	0.055	0.042	2.28 1.32 1	1.12	2.49	1.46	1.17	9.1	10.6	4.8
1×H-11	0,184	0,135	0.046	1.54 1.90 1	1.08	1.72	2.04	1.14	11.8	7.4	5.7
PXH-12	0.170	0.056	0.103	1.67 1.30 1	1.42	1.85	1.45	1.47	10.8	11.3	3.6
PXE-13	0.072	0.144	0.115	1.09 1.96 1.	.30	1.25	2.09	1,38	14.9	6.4	5.9
PXE-14	0,123	0.037	0,181	1.43 1.16 1.	80	1.57	1,26	1.93	7.6	8.5	7.1
PXH-15	0.142	0.111	0.083	1.44 1.78 1	. 24	1.58	1.88	1.30	9,8	5.6	4.4
PXH-16	0.150	88		1,38 1,86 1,	,23	1.52	1.93	1.31	7.6	7.3	0.9
EX0-9	0.071	0.216	0.050	1.06 2.62 0.	66.	1.24	2.74	1.08	16.8	4.4	8.3
FX0-10	0.165	0.113	0.045	1.62 1.96 0.	16.	1.79	2.06	1,08	10.1	5.0	11.0
FX0-11	0.254	0.041	0.049	2.59 1.30 1.	•02	2.71	1.38	1,11	4.8	6.2	8.2
FX0-12	0.168	0.044	660.0	1,90 1,36 1,	• 25	2.11	1.44	1,28	10.9	6.1	2.3
1	1				1	1	! ! !	1		1 1	

Table 5.8 (contd)

	Equilibri	Equilibrium composition in	tion in	Molecules ad	lsorbe	d per	cavity				×
Kun	liquid phe	liquid phase, mole fraction	fraction	Experimental Predicted		Pre	dicted		Per ce	Per cent, Error	1711
	-1	2	3	1 5	3	7	2	2	-	2	23
PH0-9	0.070	0.043	0.185	1.30 1.16 1.87	1,87	1.49	1.49 1.24 1.92	1.92	15.0	7.2	2.6
PH0-10	0.160	0.101	0.055	1.91 1.56 1.09	1,09	2,08	2.08 1.65 1.15	1.15	8.7	5.7	5.4
PHO-11	0,168	0.046	0.114	2,01 1,18 1,30	1.30	2.20	1.26 1.38	1.38	9.3	9,3 6,8	6.1
PH0-12	0.270	0.039	0.054	2,84 1.08 1.05	1.05	3,02	3,02 1,20 1,13	1.13	6.2	6.2 11.4	7.8
					THE R. W. P. SECOND G. P. SECOND.			-			Ì

Note: Natations are same as used in Table 5.6

adsorption model is not holding good. Many predicted values are more than 10 per cent only because all of the ternary systems studied had pentane as one of the component, for which thermal vibrations and hence errors are more at 30° C. From the comparison between predicted and experimental values for binary and ternary systems, it seems that maximum discrepancy lies in systems, at 30° C, having pentane as one of the components.

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CHAPTER 6

CONCLUSIONS AND RECOMMENDATIONS

6.1 CONCLUSIONS:

The following conclusions were made from this study:

- l. An adsorption cell was developed and used for liquid phase adsorption studies on pure and multicomponent systems. Pure, binary and ternary systems of n-pentane, n-hexane, n-heptane and n-octane, with benzene as a diluent, were studied at 6, 18, 30 and 42°C.
- 2. In adsorption of single components, lower molecular weight paraffins were preferentially adsorbed. For all the paraffins studied, adsorption decreased with increase in temperature; the decrease was largest for the lightest component.
- 3. Upto 18°C, for all the binary systems studied, temperature had no effect on the equilibrium adsorption. At 30°C, there was a decrease in adsorption of pentane for all the systems containing it as one of the components. For the systems studied this decrease was highest for pentanehexane system and the lowest for pentane-octane system. Because of benzene as a diluent, conventional separation factor was found to be concentration dependent. The separation factor was redefined to make it independent of

concentration. Except for binary systems at 30°C containing pentane, the redefined separation factor was independent of temperature. At 30°C it was found to decrease for pentane containing binaries, being the lowest for pentanehexane system.

4. A two parameter statistical model was developed to represent pure component adsorption data. This model was then extended for binary and ternary systems and was validated by experimental data. Predictions by proposed model, for all the binary and ternary systems, were in good agreement with experimental adsorption data for temperatures up to 18°C. Errors were more for the systems containing pentane at 30°C.

6.2 RECOMMENDATIONS:

The present study does not cover all the n-paraffins of industrial importance. It is, therefore, recommended to perform adsorption studies on the remaining paraffins present in petroleum fractions like naphtha, kerosene, etc. With complete data on pure systems one should be able to predict multicomponent equilibrium loadings necessary for the design of industrial separation systems. The proposed model, with no clues available for liquid phase adsorption, seems to be a reasonable first attempt. In this model

may be critical near the boiling point of a component and hence for future it is recommended to take this into account also. Since in any petroleum fraction, there are more than three components present, it is, therefore, recommended to test the model for the systems with more components.

APPENDIX A3.1

CALIBRATION OF GAS CHROMATOGRAPH

In the present study, for all the cases, benzene was used as a reference material., Standardization curves, for all the components, were obtained using known mixtures of paraffin in benzene. The peak areas were measured by a disc integrator attached to the recorder. The plot of peak area ratio of paraffin to benzene versus respective known weight ratio gave a linear standardization curve for each paraffin. A typical chromatogram is shown in Figure A3.1 and these calibration curves are shown in Figure A3.2. Operating conditions for analysis were:

Chromatograph: Chromatography and Instruments Co.

Baroda, India

Carrier gas: Nitrogen

Flow rate: 25 ml/min

Column dimensions: 1/8 inch x 30 ft, S.S.

Packing: 20 per cent bentone on Chromosorb W

Column and detector temperature: 90°C

Bridge current: 100 mA

Detector: Thermal conductivity (Katharometer)

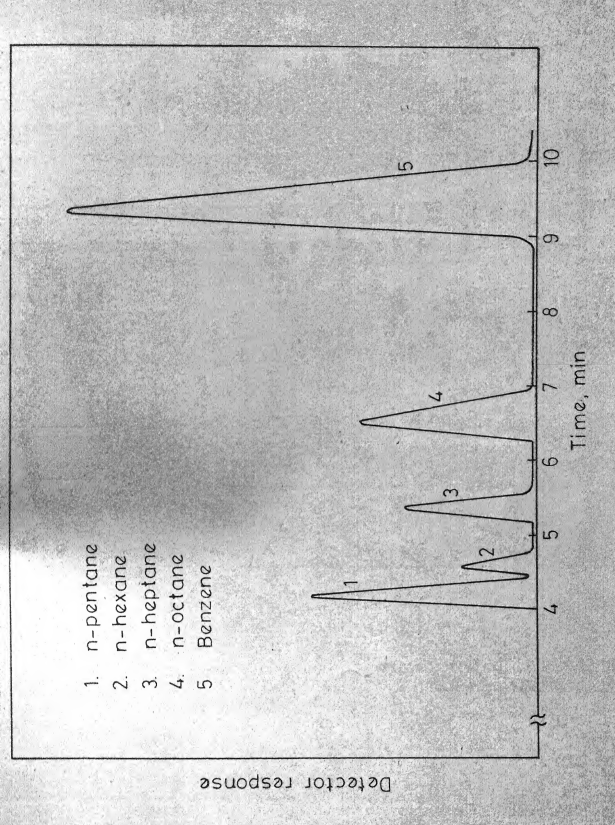


Fig.A3.1 - A typical chromatogram for n-parattins-benzene mixture.

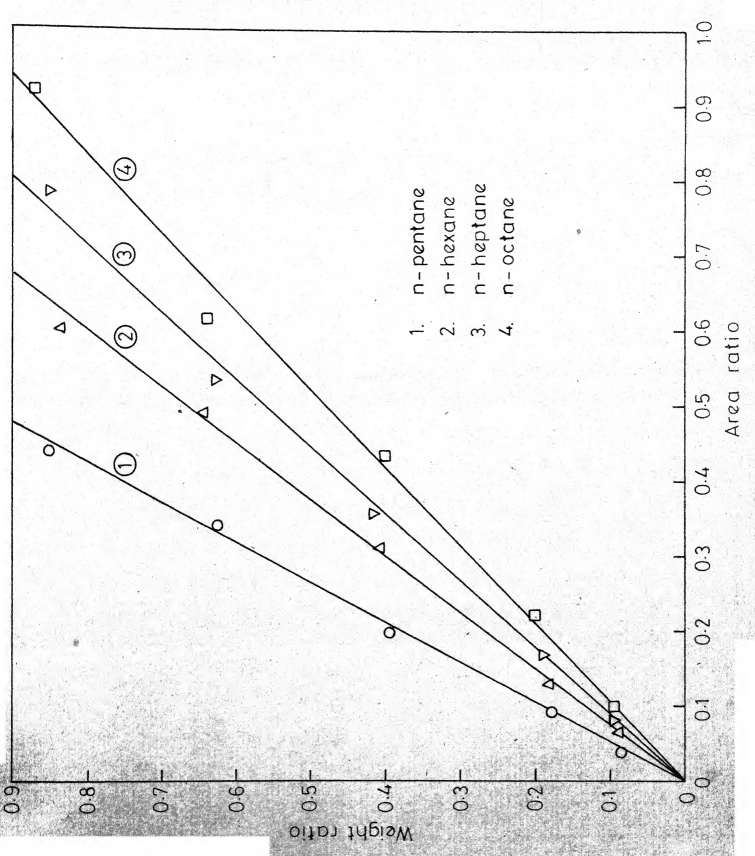


Fig. 43.2-Calibration curves for different paratfins.

APPENDIX A3.2: EXPERIMENTAL RESULTS FOR BREAKTHROUGH

CURVES OF n-HEPTANE

Technic Sections of the Control of States and Control Section Sections (Control Section Sec		Andrew Control of the	
30	O _O C		42 ⁰ C
Weight of a	dsorbent=30.1692 gms	Weight of a	dsorbent=30.2937 gms
Initial cond	centration =4.05moles/ litre	Initial con	centration = 2.35mol litre
Time, min.	Amount adsorbed, gms/gm adsorbent	Time, min.	Amount adsorbed, gms/gm.adsorbent
5	0.081	5	0.072
13	0.114	16	0.121
30	0.117	33	0.127
40	0.120	65	0.129
60	0.119	110	0.133
105	0.122	180	0.133
165	0.122	250	0.134
205	0.123	310	0.137
340	0.125	500	0.138
425	0.124	860	0.140
555	0.123	1120	0.142
675	0.127	1560	0.140
880	0.124	1650	0.141
1370	0.125	2250	0.141
1900	0.126	2950	0.142
2740	0.123	4005	0.141
3860	0.125		

APPENDIX A3.3: EXPERIMENTAL RESULTS FOR PURE COMPONENTS AT 6°C

					And strong to a first designation in particular section in a section of the secti
Run	Initial weight of paraffin, gms.	Weight of benzene, gms.	Weight of Edsorbent, gms.	Equilibrium composition in liquid phase, moles/litre	Amount adsorbed x103, moles/gm.adsorbent
PCP-1	6.2266	34.6860	28.8314	0.613	2.11
PCP-2	7.9232	32.0250	30.5984	0.957	2.34
PCP-3	9,9353	29,4294	31.1904	1.571	2.41
PCP-4	11.5071	27.3132	30.0712	2,135	2.46
PCP-5	12.2716	26.0300	29.5984	2,359	2.60
PCP-6	14.7726	22.5500	30,8231	3.134	2.69
1					
PCX-1	6.4 789	34.6949	30,3089	0.531	1.74
PCX-2	8.9915	31,2825	30.0537	1,184	1.85
PGX-3	11,1530	27.6715	30,1619	1,808	1.88
PGX-4	12,4690	25.9012	30.2414	2.177	1.90
PCX-5	14,0024	24.1504	30.1572	2,562	1.98
PCX-6	15.5839	21,5524	30.1579	3.052	2,00
1			mand make blood these trans larger		1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1

Appendix A3.3 (contd)

Run	Initialweight of paraffin, gms.	Weight of benzene, gms.	Weight of adsorbent, gms.	Equilibrium composition in liquid phase, moles/litre	Amount adsorbed x103, moles/gm.adsorbent
PCH-1	5.9552	37.0112	30.0612	0.369	1.44
FCH-2	7.3584	34.6630	30,9299	0.612	1.53
PCH-3	9,6718	31,1180	30.0182	1.159	1.61
PCH-4	13.6947	25.9943	30.1776	2.093	1.65
PCH-5	15,5407	21,6310	30.1525	2,667	1,68
PCH-6	19.2762	23,1126	29.9446	3.040	1.77
1					;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;
77	4.7286	39.4436	29,4436	0.102	1.25
PC0-2	8,6003	35,0220	29,7638	0.757	1.40
PC0-3	13,3325	30,5282	29,8819	1.598	1.46
PC0-4	18,1627	26.1594	30,1518	2.408	1,48
	-	The state of the s	TIVE		

PCH = Heptane; PCO = Octane Notes: PCP = Pentane ; PCX = heXane ;

APPENDIX A5.4: EXPERIMENTAL RESULTS FOR PURE COMPONENTS AT 180C

Run	Initial weight of paraffin, gms.	weight Weight of Ifin, benzene, gms.	Weight of adsorbent, gms.	Equilibrium composition in liquid phase, moles/litre	Amount adsorbed x 10 ³ , moles/gm.adsorbent
PCP-7	5,5164	34,2741	30,1516	0.514	1,84
PCP-8	9698.9	32,1562	29,8920	0.897	1,97
PCP-9	9.7070	28.3424	29,8155	1.805	2.06
PCP-10	11.6121	26.1195	30,4224	2,396	2,08
PCP-11	13.0938	23.8120	30,0950	2.900	2,14
PGP-12	14.6541	21,5422	30.3362	3,381	2,26
1		1 1 1 1	1 1 1 1 1 1		
PCX-7	4.8062	35,8121	30,2010	0.305	1.42
PCX-8	6.6721	33,3212	30.1327	0.738	1.55
PCX-9	8,4502	33,1627	29,7911	1,204	1.60
PCX-10	10.8467	27,0012	30.2712	1,888	1.63
PCX-11	13.6404	23.4124	30.1124	2.645	1.71
PGX-12	15.2694	21.4993	30,3313	3.056	1.77
1	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1				

Appendix A3.4 (contd)

Equilibrium composition in liquid phase, moles/litre 0.240 0.635 1.403 2.181 2.813 3.199	Weight of adsorbent, gms. 30.1224 30.0763 29.9427 30.2729 29.7539 29.7539 29.7538 29.8819	Weight of benzene, gms. 37.8911 34.6521 29.7512 25.8162 22.6412 21.5222 21.5222 21.5222 39.4436 35.0220	Initial weight Weight of paraffin, gms. 4.9563 37.8911 6.8680 34.6521 10.2145 29.7512 15.7656 25.8162 16.7123 22.6412 19.4322 21.5222 5.0870 39.4436 9.5649 35.0220 14.6110 30.5282
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Note: Notations are same as used in Appendix A3.3

APPENDIX A3.5: EXPERIMENTAL RESULTS FOR PURE COMPONENTS AT 30°C

Run	Initial weight of paraffin, gms.	weight Weight of fin, benzene, gms.	Weight of adsorbent, gns.	Equilibrium composition in liquid phase, moles/litre	Amount adsorbed x 167 moles/gm.adsorbent
PCP-13	4.9792	36.8414	30,1627	0.491	1.56
PCP-14	6,1819	35,4624	30.2117	0.774	1,69
PGP-15	7.1965	33.6217	29.9648	1.067	1.76
PCP-16	8.7734	29,8030	29,8984	1.580	1.85
PCP-17	11,3450	23.0523	30.2504	2,616	1.90
PCP-18	13,2071	20,1176	30.3348	3.298	1.95
1	the same bear here and here have a	Danie Mr. M. Second Paper Second Se			1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
PCX-13	3.9664	38,2135	30.2250	0.159	1.29
PCX-14	5.2980	36,6232	30.2590	0.476	1-33
PCX-15	6.5841	34.8473	30.1720	0.787	1.37
PCX-16	10,8198	27.6433	30,1517	1.938	1.42
PCX-17	14,4555	24.3121	30,2237	2.764	1.52
PCX-18	15.1436	20,8467	30.0917	3,126	1.60
		- 1			
	the second through the second th				

Appendix A3.5 (contd)

Run	Initial weight Weight of paraffin, benzeners.	Weight of benzene, gms.	Weight of adsorbent, gms.	Equilibrium composition in liquid phase, moles/litre	Amount adsorbed z moles/gm.adsorbent
PCH-13	4.1599	38.6491	29.6542	0.129	1.20
PCH-14	5.4954	36.4382	30.1355	0.384	1.26
PCH-15	6.9523	34.7820	29,1270	0.713	1,29
PCH-16	9,6658	31,2298	29.8075	1.290	1.32
PCH-17	15.0776	23.1127	30.2279	2,564	1.34
PCH-18	19,2518	20,1729	30,1617	3.280	1.46
1 1			peed pack spine term many seems		
PG0-9	5.0126	39,4436	29.8127	0,205	1,15
PC0-10	9.4363	35.0220	29,7638	0.938	1,28
PC0-11	14.1260	30,5282	29.8819	1.734	1,29
PC0-12	19,7864	26,1594	30,1518	2,571	1,31
		Annual Resident Market State College of the College			

Notations are same as used in Appendix AJ. J Note:

APPENDIX A3.6: EXPERIMENTAL RESULTS FOR PURE COMPONENTS AT 42°C

Run	Initial weight of paraffin, gms.	Weight of benzene, gms.	Weight of adsorbent, gms.	Equilibrium composition in liquid phase, moles/litre	Amount adsorbed $x 10^3$ moles/gm. adsorbent
PCX-19	3,9565	38.4122	30.1729	0.173	1,26
PGX-20	5.2359	36.8409	29,7009	0.488	1.29
PGX-21	7.9588	33.1919	30,3013	1.144	1.32
PGX-22	10.5224	29,9335	30,6614	1.771	1.34
PGX-23	14.7546	25.0412	30.0920	2.771	1.40
PCX-24	16,3813	22,4673	29,8512	3,188	1.47
PCH-19	4.4595	38.5281	30.2214	0.183	1,19
PCH-20	5.6869	36,3121	29,8729	0.444	1.23
PCH-21	7.8807	33.6171	29,9253	0.908	1.25
PCH-22	13.5459	26,1927	30,0625	2,142	1.29
PCH-23	18,0898	23,7676	30,3016	2,873	1.33
PCH-24	22.0978	21,0921	30,2842	3,451	1.45
PC0-13	5.0558	39.4436	29.8127	0.236	1.11
PC0-14	9.9295	35.0220	29,7638	1,022	1.23
PC0-15	14.4930	30,5282	29,8819	1.770	1,25
PC0-16	20.6542	26.1594	30,1518	2,626	1.27
Note: No	Notations are same	as used in	Appendix A3	3	130

Appendix A4.1 (contd)

Run	Initial weight of paraffins, gms.	0 H	Weight of benzene, gms.	Weight of adsorbent, gms.	Equilibrium composition in liquid phase, mole fracti	quilibrium compo- sition in liquid phase, mole fraction	Adsorbe compos mole fr	Adsorbed phase composition; mole fraction .	Separation factor, K
BPH-1	14.1358	3.3475	21,7080	30,3614	0.877	0.123	0,815	0.185	0.62
BPH-2	10.9897	5.9411	21,7019	30.5584	0.716	0.284	0.727	0.273	1,06
BPH-3	7,2624	9.6524	21,6857	30.8941	0.510	0.490	0.554	0.446	1,29
BPH-4	3.3264	15.0246	21.6963	29,5329	0.198	0.802	0.318	0.682	1.88
BP0-1	2.8074	17.5999	21.7013	30.2394	0.165	0,835	0.290	0.710	2,06
BP0-2	6.2894	14.7825	21.6429	30,1516	0.409	0.591	0.506	0.494	1.48
BP0-3	9.7731	8.0076	21.7214	29,8294	0.653	0.347	0.673	0.327	1,09
BP0-4	14,5298	4.3224	21.7398	29.7286	0.857	0.143	0,811	0.189	0.72
		-	The second secon		The state of the s				

anhese mole fractions are on benzene free basis,

Note: 1 represents the lighter hydrocarbon in a given binary mixture and 2 presents the other component.

BPH = Pentane-Heptane BHO = Heptane-Octane; BXH = heXane-Heptane; BPX = Pentane-heXane; BPO = Pentane-Octane

APPENDIX A4.2: EXPERIMENTAL RESULTS FOR BINARY SYSTEMS AT 18°C

manse. benzene, gms. adsorbent, tion in liquid phase, composition, and extraction in liquid phase, composition, and extraction in liquid phase, and extraction in liquid phase, composition, and extraction, and extraction in liquid phase, composition, and extraction,	Initial w	weight of	Weight of	Weight of	11.6	COD	Adsorbed	ed phase	ti
21.6802 30.0021 0.865 0.137 0.602 0.209 21.6915 30.1527 0.660 0.340 0.662 0.338 21.6929 29.9284 0.428 0.572 0.472 0.528 21.6815 30.2721 0.182 0.818 0.283 0.717 21.6902 29.8827 0.186 0.814 0.387 0.613 21.6902 29.7625 0.421 0.579 0.492 0.508 21.6992 30.1029 0.831 0.169 0.736 0.264 21.6826 29.8816 0.174 0.826 0.518 0.562 21.6825 29.9222 0.407 0.593 0.458 0.542 21.7127 30.1713 0.622 0.378 0.589 0.411 21.7127 30.1712 0.854 0.723 0.723 0.277	paraffins, 1	em s. 2	benzene, gms.	adsorbent, gms.	jr. Le	phase	compc mole l	ition, raction 2	Factor, K
21.6915 30.1527 0.660 0.340 0.662 0.358 21.6929 29.9284 0.428 0.572 0.472 0.528 21.6815 30.2721 0.182 0.818 0.283 0.717 21.6613 29.8827 0.186 0.814 0.587 0.613 21.6902 29.7625 0.421 0.579 0.492 0.508 21.6992 30.1029 0.831 0.169 0.736 0.264 21.6364 29.8816 0.174 0.826 0.318 0.682 21.6825 29.9222 0.407 0.593 0.458 0.542 21.7127 30.1713 0.622 0.378 0.589 0.411 21.7038 29.7727 0.854 0.146 0.723 0.277		3.2372	21,6802	30,0021	0.863	0.137	0.791	0.209	09*0
21,6929 29,9284 0.428 0.572 0.412 0.528 21,6815 30,2721 0.182 0.818 0.283 0.717 21,6613 29,8827 0.186 0.814 0.387 0.613 21,6902 29,7625 0.421 0.579 0.492 0.508 21,692 30,1895 0.628 0.372 0.632 0.368 21,692 30,1029 0.831 0.169 0.736 0.264		6961.9	21.6915	30.1527	099*0	0.340	0.662	0.338	1.01
21,6815 30,2721 0,182 0,818 0,283 0,717 21,6613 29,8827 0,186 0,814 0,387 0,613 21,6902 29,7625 0,421 0,579 0,492 0,508 21,6992 30,1029 0,831 0,169 0,736 0,264 21,6992 30,1029 0,831 0,169 0,736 0,264 21,6364 29,8816 0,174 0,826 0,318 0,682 21,6825 29,9222 0,407 0,593 0,458 0,542 21,7127 30,1713 0,622 0,378 0,589 0,411 21,7038 29,7727 0,854 0,146 0,723 0,277	-	10,1203	21.6929	29,9284	0.428	0.572	0.472	0.528	1,19
21.6613 29.8827 0.186 0.814 0.587 0.613 21.6902 29.7625 0.421 0.579 0.492 0.508 21.7107 30.1895 0.628 0.372 0.632 0.568 21.6992 30.1029 0.831 0.169 0.736 0.264 21.6364 29.8816 0.174 0.826 0.318 0.682 21.6825 29.9222 0.407 0.593 0.458 0.542 21.7127 30.1713 0.622 0.378 0.589 0.411 21.7038 29.7727 0.854 0.146 0.723 0.277			21,6815	30.2721	0.182	0.818	0.283	0.717	1.78
21.6902 29.7625 0.421 0.579 0.492 0.508 21.7107 30.1895 0.628 0.372 0.632 0.368 21.6992 30.1029 0.831 0.169 0.736 0.264			21.6613	•	0.186	0.814	•	0.613	2.76
21.7107 30.1895 0.628 0.372 0.632 0.368 1. 21.6992 30.1029 0.831 0.169 0.736 0.264 0. 21.6364 29.8816 0.174 0.826 0.318 0.682 2. 21.6825 29.9222 0.407 0.593 0.458 0.542 1. 21.7127 30.1713 0.622 0.378 0.589 0.411 0. 21.7038 29.7727 0.854 0.146 0.723 0.277 0.		11,1023	21.6902	29,7625	0.421	0.579	0.492	0.508	1.33
21.6992 30.1029 0.831 0.169 0.736 0.264 0. 21.6364 29.8816 0.174 0.826 0.318 0.682 2. 21.6825 29.9222 0.407 0.593 0.458 0.542 1. 21.7127 30.1713 0.622 0.378 0.589 0.411 0. 21.7038 29.7727 0.854 0.146 0.723 0.277 0.		7.5245	21.7107	30,1895	0.628	0.372	0.632	0.368	1,02
21.6364 29.8816 0.174 0.826 0.518 0.682 2. 21.6825 29.9222 0.407 0.593 0.458 0.542 1. 21.7127 30.1713 0.622 0.378 0.589 0.411 0. 21.7038 29.7727 0.854 0.146 0.723 0.277 0.		4,2015	21.6992	30,1029	0.831	0,169	0.736	0.264	0.56
21.6364 29.8816 0.174 0.826 0.318 0.682 2. 21.6825 29.9222 0.407 0.593 0.458 0.542 1. 21.7127 30.1713 0.622 0.378 0.589 0.411 0. 21.7038 29.7727 0.854 0.146 0.723 0.277 0.	1	-1			1				
21.6825 29.922 0.407 0.593 0.458 0.542 21.7127 30.1713 0.622 0.378 0.589 0.411 21.7038 29.7727 0.854 0.146 0.723 0.277		15.0288	21,6364	29,8816	0.174	0.826	0,318	0.682	2.22
21.7127 30.1713 0.622 0.378 0.589 0.411 21.7038 29.7727 0.854 0.146 0.723 0.277		10.1608	21,6825	29,9222	0,407	0.593	0.458	0.542	1,23
21.7038 29.7727 0.854 0.146 0.723 0.277		7.0137	21,7127	.17	0.622	0.378	. 58	0.411	78.0
		3.7221	21,7038	.77	0.854	0.146	•72	0.277	0.44

Appendix A4.2 (contd)

					Control of the Agency part designed by the Agency by the A				
Run		Initial weight of paraffins, gms.	Weight of benzene, gms.	Weight of adsorbent, gms.	Equilibri tion in 1 mole frac	Equilibrium composition in liquid phase, mole fraction 2	Adsorbe composi mole fr	Adsorbed phase composition, mole fraction	Separation factor, K
BPH-5	14.1038	3 4302	7083 [6	7,00,00			to the same of the	And the control of th	
			7000.47	1700.67	9/8.0	0.124	0.793	0.207	0.54
BPH-6	10.6794	5.9672	21,6892	30,1516	0.712	0.288	0.716	0.284	1,02
BPH-7	7.7114	9.5931	21.6712	30.2910	0.513	0.487	0.565	0.435	20.1
BPH-8	3,3628	14.5987	21,6698	30,0152	0.209	0,791	0.329	0.671	1.85
1 1 1	I I I I	Manage Property Control Contro	A American secure secure secure	1	1	; ; ;			\ !
BP0-5	3,2892	15.4744	21,7185	29.8326	0.212	0.788	0.349	0.651	1 1 00 1
BP0-6	6.8179	10.7777	21,6623	29,8129	0.479	0.521	0.554	0.446	ן ר ט א
BP0-7	10.2713	7.6230	21,6629	29,5027	0.675	0.325	969*0	0.304	
BP0-8	14.5588	3.8900	21,6903	30.089	0.879	0.121	0,802	0.198	1 U
								3	0

agnese mole fractions are on benzene free basis Note: Notations are same as used in Appendix A4.1

APPENDIX A4.3: EXPERIMENTAL RESULTS FOR BINARY SYSTEMS AT 30°C

Separation factor, K	1	1.44	0.72	0.49	0.34		0.46	1.10	1.56	2.57	I I I I	2,73	1.50	0.95	0.45	
bed phase sition, fraction	2	0.759	0.627	905.0	0.311	1 1 1	0.255	0.329	0.439	0.623	1 1 1	0.678	0.558	0.412	0.294	
Adsorbed ph composition mole fracti	Н	0.241.	0.373	0.494	689.0	1	0.745	0.671	0.561	0.377	1 1	0.322	0.442	0.588	901.0	-
um compo- liquid e fraction a	2	0.795	0.547	0.333	0.134	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	0.137	0.350	0.550	0.810		0.852	0.655	0.399	0.157	
Equilibrium sition in l phase, mole	1	0.205	0.453	199.0	998.0	1	0.863	0.650	0.450	0.190	1	0.148	0.345	0.601	0.843	
Weight of adsorbent, gms.		29.8627	30,1690	30.0854	30.0989	1 1 1 1	29,8314	29,7729	29,9238	30,1122	1 1 1 1	29,7622	29,8320	29,9698	29,7440	
Weight of benzene, gms.		21.5863	21,8855	21.9107	21,9169	1	21.7025	21,6817	21,6939	21,6898	1 1 1	21.6396	21,7012	21,6988	21,6628	
weight of ns,ems. 2		13.9764	10.6777	6.9730	3.6111		3.6045	7.7851	10,1209	14.5098	1 1 1	16,7999	11.7229	7.4713	3.9464	
Initial weight o paraffins, gms.		3,3295	9898.9	9.7702	13.9554		15.4862	12,6950	7,9399	3.8784	1 1		6.0619	9.7249	14.4074	
Run		BPX-9	BPX-10	BPX-11	BFX-12		BXH-9	BXH-10	BXH-11	BXH-12		1	BH0-10	BH0-11	BH0-12	

Appendix A4.3 (contd)

Run	Initial weight of paraffins, gms.	ight of 18, 2	Weight of benzene, gms.	Weight of adsorbent, gms.	Equilibr sition i phase, m	Equilibrium composition in liquid phase, mole fraction l	Adsorbed phase composition, mole fraction 1	Adsorbed phase composition, mole fraction.	Separation factor, K
BPH-9	14.3499 3.7235	3.7235	21.6517	30.1516	0.877	0.123	0.744	0.256	0.41
BPH-10	11,1221 (6.5071	21,6899	30,2912	0.725	0.275	0.643	0.357	0.68
BPH-11	8.0645 10.0740	0.0740	21.6712	30.0459	0.534	0.466	0.501	0.499	0.88
BPH-12	3,4456 14,6269	4.6269	21.6025	30.1127	0.220	0.780	0.279	0.721	1.66
		1			1	NOT NAME AND POST OFFICE AND POST			
6-0	3,4364	17.3266	21,6626	30.1124	0.204	962.0	0.346	0.654	2.07
BF0-10		12,6457	21.6439	29,8826	0.465	0.535	0.539	0.461	1,35
BP0-11	10.7569	8.0406	21,6827	29,8412	619.0	0.321	619.0	0.321	I.00
BP0-12	14.3044	4.2407	21,6836	29,9018	998.0	0.134	0.781	0.219	0.55

agnese mole fractions are on benzene free basis

Note: Notations are same as used in Appendix A4.1

ARPENDIX A4.1: EXPERIMENTAL RESULTS FOR BINARY SYSTEMS AT 42°C

Run	Initial weight paraffins, gms.	Initial weight of paraffins, gms.	Weight of benzene,gms	Weight of adsorbent, gms.	Equilibrium composition in liquid phase, mole fractio	m compo- liquid fraction 2	Adsorbed phase composition, mole fraction l	Separation factor, K
BXH-13 3.6740	3.6740	13.9992	21.6717	29,8010	0.189	0.811	0.378 0.622	2,61
BXH-14	7.4603	10,0817	21,6829	29,7700	0.434	0.566	0.555 0.445	1.62
BXH-15 11,4451	11,4451	6.7269	21,6528	30,1217	0.662	0.338	0.670 0.330	1.03
BXH-16	BXH-16 14.6661	3.5147	21,7512	30.0129	0,860	0.140	0.733 0.267	0.45
IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII			beine these parts taken give and	teres been tento pued freque				* *** *** *** *** ***
BH0-13	BHO-13 3.5617	16.4160	21.6538	30,1616	0.154	0.846	0.328 0.672	2,68
BH0-14	6.9718	13.5424	21,7216	29,8517	0.349	0.651	0.440 0.560	1.47
BH0-15	BHO-15 10,2128	7.5678	21,6982	29,9114	0.613	0.387	0.587 0.413	06*0
91-0ня	ыно-16 14.4486	3.9645	21,6724	29.8434	0.844	0.156	0,701 0,299	0.43
		and the second s		The state of the s	The second secon	The state of the s		

anhese mole fractions are on benzene free basis

Note: Notations are same as used in Appendix A4.1

APPENDIX A4.5: OPERATING CONDITIONS FOR TERNARY SYSTEMS AT 6°C

Run	Initial weight	of 5	paraffins, gms 3	Weight of benzene, gms.	Weight of adsorbent, gms.
PXH-1	10,6388	3.2586	5.3818	21,6916	30.2192
PXH-2	3.9274	9.2417	5.5638	21,6836	30,1814
PXH-3	2,8052	7.4966	9.4497	21,6876	29,9384
PXH-4	9951-9	5.1188	8,3040	21.6746	0
PX0-1	9.2721	3.0436	3.8059	21.6265	29,8626
PX0-2	3.8967	7,8161	3,7385	21,6629	29,8884
PX0-3	2.7226	6.9346	5.9805	21,6778	29,9254
PX0-4	5.6098	4.9040	6.1882	21,6828	30.1146
PH0-1	10.1738	3.8783	4.1975	21,7108	29,9827
PH0-2	3.8141	5.1505	6,1502	21,6927	30.0839
PH0-3	2,6733	9.0024	3.8486	21,6634	30,2256
PH0-4	2,8028	7.3864	5.5603	21,706.8	30,1617
Note:	1 is the lightest hyd hydrocarbon.		rocarbon in a gi	given ternary system and	3 is the heaviest
PXH =	Pentane-heXane-Heptane	Heptane;	PXO = Penta	Pentane-heXane-Octane;	138
DHO =	Dontone Henters	Octobor			

Note: Notations are same as used in Appendix A4.5

APPENDIX A4.6: OPERATING CONDITIONS FOR TERNARY SYSTEMS AT 18°C

		× ×	Committee of the Commit		en aghendaran bendarangan penancangan penancangan diberapat aban penancangan dan bendaran anakan menancanan me
Run	Initial weight 1	of	paraffins, gms 3	Weight of benzene, gms.	Weight of adsorben gms
PXH-5	5.0132	3.1772	9,3806	21.6713	30.1256
PXH-6	5.2235	6.3778	6.4338	. 21.6527	29.7943
PXH-7	9.4934	9996.9	2.5716	21.7016	29,9004
PXH-8	5.6149	8.3080	2,5063	21,6884	29,8715
1 1 1				1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	
PX0-5	2,4187	2.9722	10.2961	21.6298	29,8989
PX0-6	9.5309	4.0827	2,6465	21.6325	29.8725
PX0-7	4.9743	7.2392	3.4290	21,6386	29,9206
PX0-8	2.7661	7.5776	5,9953	21.6512	29,8127
PHO-5	4.4916	3.1085	9500.6	21.6938	29,9635
PH0-6	9.3936	3.2041	2,6660	21,6826	30.1217
PH0-7	3.0125	6,0269	6,1011	. 21,7002	30,2435
PH0-8	7.1677	5.9279	3,2853	21,6888	30.0628
				AND THE PROPERTY OF THE PROPER	

APPENDIX A4.7: OPERATING CONDITIONS FOR TERNARY SYSTEMS AT 30°C

1 2 3 mergin of gans. 1 2 3 mergin of gans. 1 2 2 3 mergin of gans. 1 2.12.0 1.2.12.0 1.2.12.0 1.2.12.0 1.2.12.0 1.2.12.0 1.2.12.0 1.2.12.0 1.2.12.0 1.2.12.0 1.2.12.0 1.2.12.0 1.2.12.0 1.2.12.1 1.2.12.2 1.2.12.0 1.2.12.	D	Initial weight of	ight of para	affins, gns.	15 to 15	0.4° 40° 40°	
5.6674 11.8669 5.3120 21.8955 0 12.1527 3.5771 3.2395 21.8900 1 8.5445 7.5322 3.5182 21.9500 2 7.9277 3.6292 6.2008 21.9147 3 3.5245 7.9038 6.7744 21.8947 4 5.9696 2.7866 10.4366 21.8929 5 6.5736 6.2677 5.2552 21.8929 6 7.1555 7.7968 6.4614 21.8749 7 10.2570 3.9158 21.7015 1 10.2570 3.0167 4.5387 21.6998 2 6.7440 3.1244 6.4821 21.6986	Kun	Т	2	2	ougram	or penzene, gms.	werght of ausorbemt, gms.
12.1527 3.2395 21.8900 8.5445 7.5332 3.5182 21.9500 7.9277 3.6292 6.2008 21.9147 5.9696 2.7866 10.4366 21.8947 6.5736 6.2677 5.2552 21.8929 7.1555 7.7968 6.4614 21.8749	PXH-9	3.6674	11,8669	3.3120		21,8955	30,2529
8.5445 7.5332 3.5182 21.9500 7.9277 3.6292 6.2008 21.9147 5.9696 2.7866 10.4366 21.8947 6.5736 6.2677 5.2552 21.8929 7.1555 7.7968 6.4614 21.8749	PXH-10	12,1527	3.5771	3.2395		21,8900	29.9740
7.9277 3.6292 6.2008 21.9147 3.5245 7.9038 6.7744 21.8947 5.9696 2.7866 10.4366 21.8869 6.5736 6.2677 5.2552 21.8929 7.1555 7.7968 6.4614 21.8749	PXH-11	8.5445	7.5332	5.5182		21,9500	30.2944
5.9696 2.7866 10.4366 21,8947 6.5736 6.2677 5.2552 21,8929 7,1555 7.7968 6.4614 21.8749 5.1831 10.8408 3,9158 21,7015 6.4726 6.2920 3.6184 21.6927 10.2570 3.1244 6.4821 21.6886	PXH-12	7.9277	3.6292	6,2008		21,9147	29,6884
5.9696 2.7866 10.4366 21.8869 6.5736 6.2677 5.2552 21.8929 7.1555 7.7968 6.4614 21.8749 7.1555 7.7968 6.4614 21.7015 6.4726 6.2920 3.6184 21.6927 10.2570 3.0167 4.5387 21.6998 6.7440 3.1244 6.4821 21.6886	PXH-13	3.5245	7.9038	6.7744		21,8947	30,0294
6.5736 6.2677 5.2552 21.8929 7.1555 7.7968 6.4614 21.8749 5.1851 10.8408 3.9158 21.7015 6.4726 6.2920 3.6184 21.6927 10.2570 3.0167 4.5387 21.6998 6.7440 3.1244 6.4821 21.6886	PXH-14	9696*5	2,7866	10.4366		21,8869	50.2539
7.1555 7.7968 6.4614 21.8749 5.1851 10.8408 3.9158 21.7015 6.4726 6.2920 3.6184 21.6927 10.2570 3.0167 4.5387 21.6998 6.7440 3.1244 6.4821 21.6886	PXH-15	6.5736	6.2677	5.2552		21.8929	30.28.14
3.1831 10.8408 3.9158 21.7015 6.4726 6.2920 3.6184 21.6927 10.2570 3.0167 4.5387 21.6998 6.7440 3.1244 6.4821 21.6886	PXH-16	7,1555	7.7968	6,4614		21.8749	29.9409
6.4726 6.2920 3.6184 21.6927 10.2570 3.0167 4.5387 21.6998 6.7440 3.1244 6.4821 21.6886	PX0-9	3.1831	10.8408	3,9158	 	21.7015	29,9297
10.2570 3.0167 4.5387 21.6998 6.7440 3.1244 6.4821 21.6886	PX0-10	6.4726	6,2920	3,6184		21.6927	29,9765
6.7440 3.1244 6.4821 21.6886	PX0-11	10.2570	3,0167	4.5387	*	21,6998	29.8965
	PX0-12	6.7440	3.1244	6.4821		21.6886	30,1028

Appendix A4.7 (contd)

117777	ntial weigh	it of para	Intial weight of paraffins, gms	Weight of henzene	14 0 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4
	H	N	W	ems sms	gms.
PHO-9	3.2666	3.2886	11.2380	21,6813	30,2216
PH0-10	6,6120	6.2257	4.2156	21,6602	29,9107
PHO-11	8796.9	3.5144	7.3890	21.6912	29.8635
PH0-12	11.2332	3,1689	4.2958	21,7002	29.9234

Note: Notations are same as used in Appendix, A4.5

ADDENDUM

somewhat inappropriately used in the text. It is usually meant to represent the time required for the first appearance of a species from the exit of an adsorption column. In this work, it has been used to describe the time required for the attainment of adsorption equilibrium.